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60/465,844	25 April 2003 (25.04.2003)	US	60/513,975	24 October 2003 (24.10.2003)
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[Continued on next page]

(54) Title: PHOSPHONATE COMPOUNDS HAVING IMMUNO-MODULATORY ACTIVITY

(57) Abstract: The invention is related to phosphonate substituted compounds having immuno-modulatory activity, compositions containing such compounds, and therapeutic methods that include the administration of such compounds, as well as to processes and intermediates useful for preparing such compounds.

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(81) **Designated States** (unless otherwise indicated, for every kind of national protection available): AE, AG, AL, AM, AT, AU, AZ, BA, BB, BG, BR, BW, BY, BZ, CA, CH, CN, CO, CR, CU, CZ, DE, DK, DM, DZ, EC, EE, EG, ES, FI, GB, GD, GE, GH, GM, HR, HU, ID, IL, IN, IS, JP, KE, KG, KP, KR, KZ, LC, LK, LR, LS, LT, LU, LV, MA, MD, MG, MK, MN, MW, MX, MZ, NA, NI, NO, NZ, OM, PG, PH, PL, PT, RO, RU, SC, SD, SE, SG, SK, SL, SY, TJ, TM, TN, TR, TT, TZ, UA, UG, US, UZ, VC, VN, YU, ZA, ZM, ZW.

(84) **Designated States** (unless otherwise indicated, for every kind of regional protection available): ARIPO (BW, GH, GM, KE, LS, MW, MZ, NA, SD, SL, SZ, TZ, UG, ZM, ZW), Eurasian (AM, AZ, BY, KG, KZ, MD, RU, TJ, TM), European (AT, BE, BG, CH, CY, CZ, DE, DK, EE, ES, FI, FR, GB, GR, HU, IE, IT, LU, MC, NL, PL, PT, RO, SE, SI, SK, TR), OAPI (BF, BJ, CF, CG, CI, CM, GA, GN, GQ, GW, ML, MR, NE, SN, TD, TG).

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For two-letter codes and other abbreviations, refer to the "Guidance Notes on Codes and Abbreviations" appearing at the beginning of each regular issue of the PCT Gazette.

PHOSPHONATE COMPOUNDS HAVING
IMMUNO-MODULATORY ACTIVITY

This non-provisional application claims the benefit of priority under 35
5 U.S.C. § 119(e) from U.S. Provisional Patent Application Serial Nos.
60/465424, 60/465373, 60/465420, 60/465380, 60/465433, 60/465481,
60/465377, 60/465581, 60/465532, 60/465844, 60/465531, and 60/465574, all
filed April 25, 2003; and to U.S. Provisional Patent Application Serial Nos.
60/493303, 60/493310, 60/493309, and 60/493302, all filed August 7, 2003; and
10 to U.S. Provisional Patent Application Serial Nos. 60/495533, 60/495529,
60/495455, 60/495537, 60/495456, 60/495398, 60/495425, 60/495427,
60/495661, 60/495393, 60/495416, 60/495614, and 60/495417, all filed August
15, 2003; and to U.S. Provisional Patent Application Serial Nos. 60/514054,
60/513971, 60/514394, 60/513975, 60/514453, 60/514202, 60/513948,
15 60/514424, 60/514280, 60/514144, 60/513979, 60/514075, 60/513946,
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60/514021, and 60/514298, all filed October 24, 2003; and to U.S. Provisional
20 Patent Application Serial Nos. 60/532230, 60/531960, 60/532160, 60/531940,
and 60/531932, all filed December 1, 2003; and to U.S. Provisional Patent
Application Serial No. 60/532591, filed December 23, 2003; and to U.S.
Provisional Patent Application Serial No. 60/536005, filed January 12, 2004.
The entirety of all Provisional Applications listed above are incorporated herein
25 by reference.

FIELD OF THE INVENTION

The invention relates generally to compounds with immuno-modulatory
(e.g., immunosuppressant) activity.

BACKGROUND OF THE INVENTION

Improving the delivery of drugs and other agents to target cells and tissues has been the focus of considerable research for many years. Though many attempts have been made to develop effective methods for importing
5 biologically active molecules into cells, both *in vivo* and *in vitro*, none has proved to be entirely satisfactory. Optimizing the association of the inhibitory drug with its intracellular target, while minimizing intercellular redistribution of the drug, e.g., to neighboring cells, is often difficult or inefficient.

Most agents currently administered to a patient parenterally are not
10 targeted, resulting in systemic delivery of the agent to cells and tissues of the body where it is unnecessary, and often undesirable. This may result in adverse drug side effects, and often limits the dose of a drug (e.g., glucocorticoids and other anti-inflammatory drugs) that can be administered. By comparison, although oral administration of drugs is generally recognized as a convenient and
15 economical method of administration, oral administration can result in either (a) uptake of the drug through the cellular and tissue barriers, e.g., blood/brain, epithelial, cell membrane, resulting in undesirable systemic distribution, or (b) temporary residence of the drug within the gastrointestinal tract. Accordingly, a major goal has been to develop methods for specifically targeting agents to cells
20 and tissues. Benefits of such treatment includes avoiding the general physiological effects of inappropriate delivery of such agents to other cells and tissues, such as uninfected cells.

Autoimmune diseases and transplantation rejection remain major public health problems worldwide. Although drugs with immunosuppressive activity
25 are in wide use and have shown effectiveness, their clinical usefulness has been limited due to their toxicity and other side effects. Currently there is a need for new immunosuppressant agents, i.e. drugs, having improved immunosuppressant activity and pharmacokinetic properties, improved oral bioavailability, greater potency, and extended effective half-life *in vivo*. New immunosuppressant
30 agents should have fewer side effects, less complicated dosing schedules, and/or be orally active.

SUMMARY OF THE INVENTION

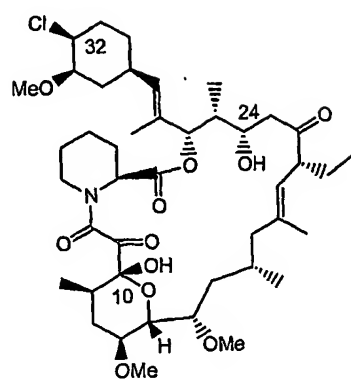
Intracellular targeting may be achieved by methods and compositions that allow accumulation or retention of biologically active agents inside cells. The present invention provides novel phosphonate containing analogs of immuno-modulatory (*e.g.*, immunosuppressant) compounds. These compounds possess the utilities of the related immuno-modulatory compounds, but due to the presence of the phosphonate group(s) they typically provide cellular accumulation of the phosphonate compound. Thus, compounds of the invention may demonstrate improved immuno-modulatory properties, pharmacokinetic properties, oral bioavailability, potency, or extended effective half-life *in vivo*, or a combination thereof. The compounds of the invention may also have distinct resistance profiles, fewer side effects, less complicated dosing schedules, or have increased oral activity.

The present invention relates generally to the accumulation or retention of therapeutic compounds inside cells. The invention is more particularly related to attaining high concentrations of phosphonate-containing molecules in target cells. Such effective targeting may be applicable to a variety of therapeutic formulations and procedures.

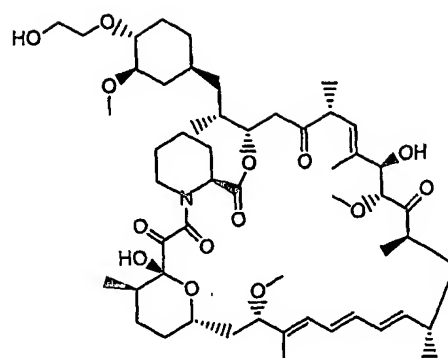
Accordingly, in one embodiment the invention provides a compound of the invention which is a conjugate comprising an immuno-modulatory compound (*e.g.*, an immunosuppressant compound) linked to one or more phosphonate groups.

Compositions of the invention include immuno-modulatory compounds having at least one phosphonate group. Accordingly, in one embodiment the invention provides a conjugate comprising an immuno-modulatory compound (*e.g.*, an immunosuppressant compound) linked to one or more phosphonate groups or a pharmaceutically acceptable salt thereof.

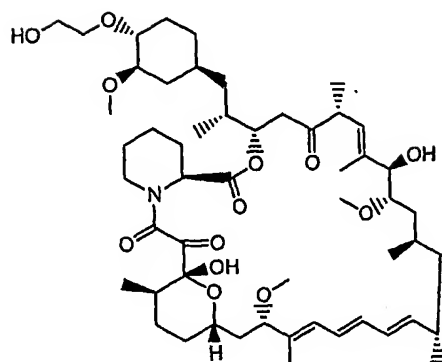
In another embodiment the invention provides a compound of any one of formulae 500-547:



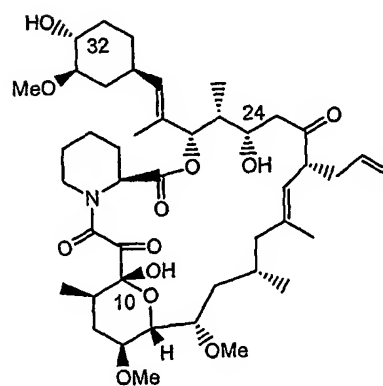
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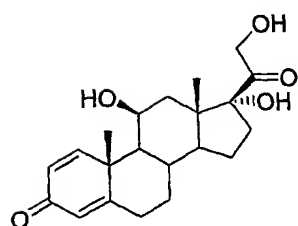


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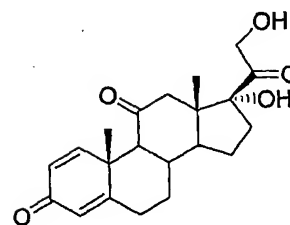


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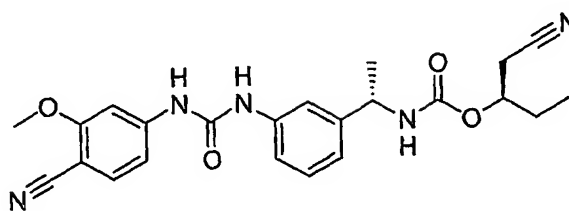
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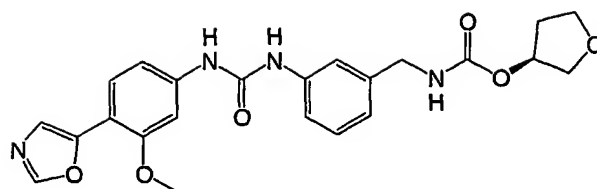


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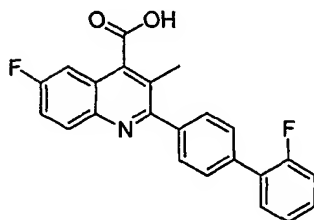


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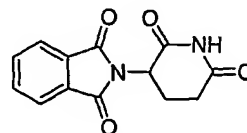
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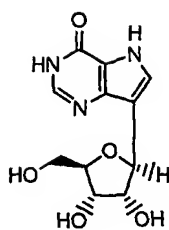
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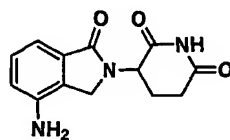
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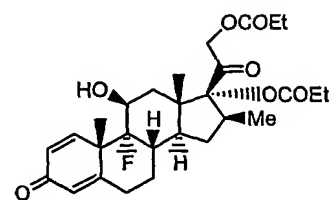
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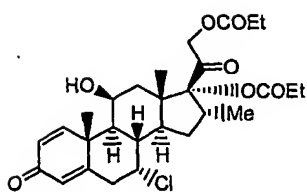
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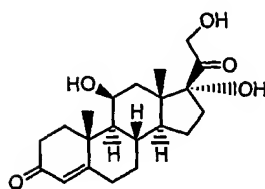
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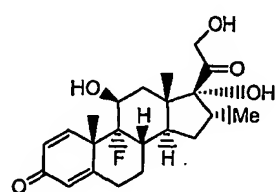
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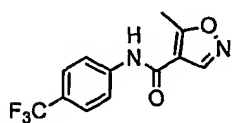
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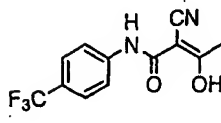
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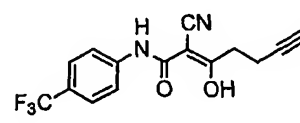
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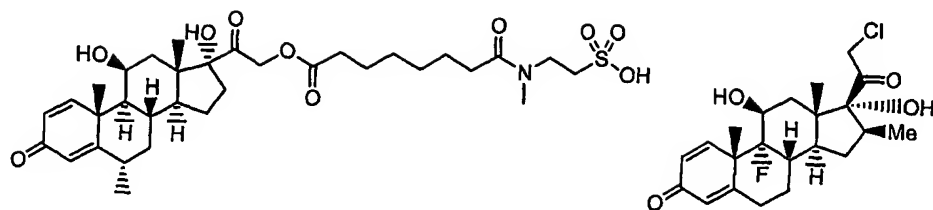
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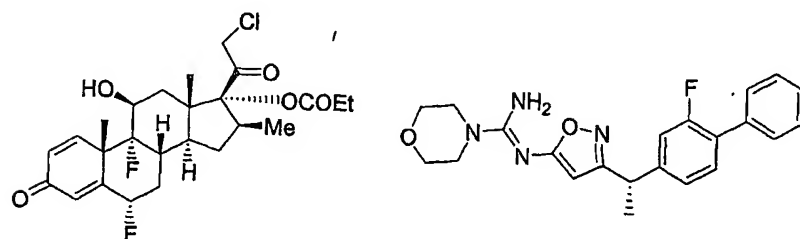


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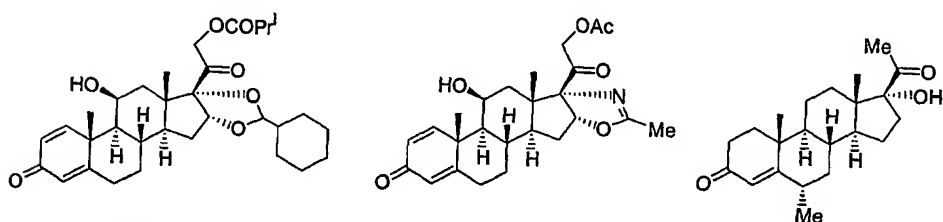
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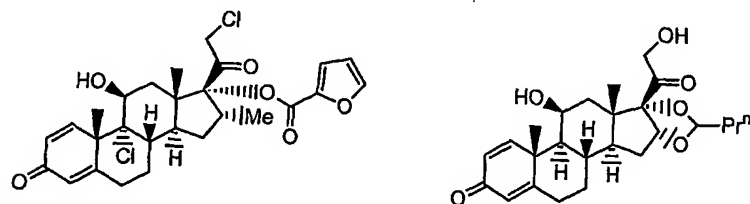
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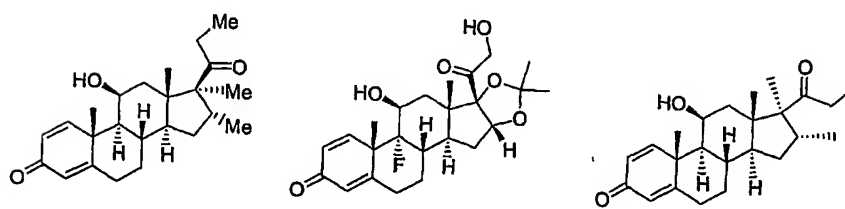
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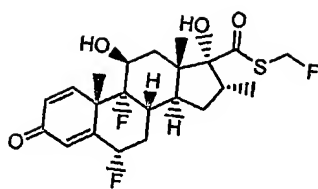
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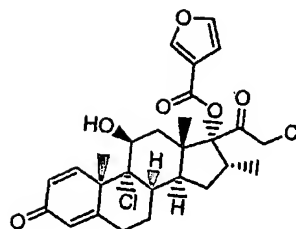
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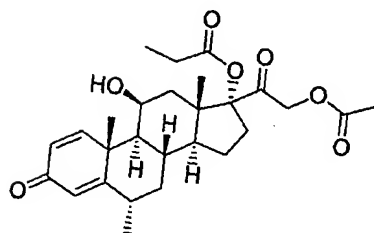
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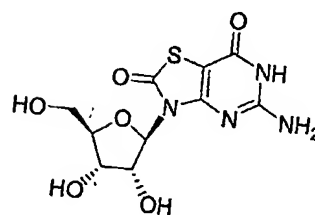
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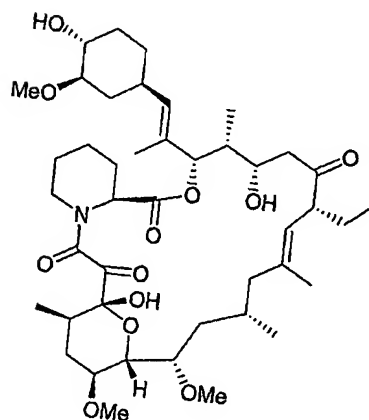
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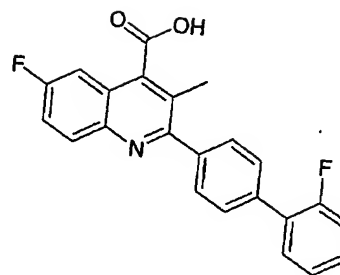
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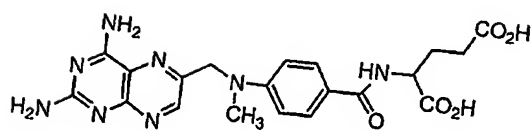
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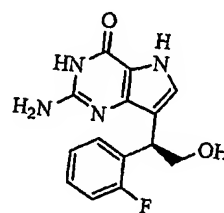
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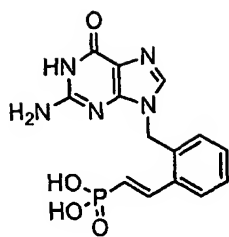
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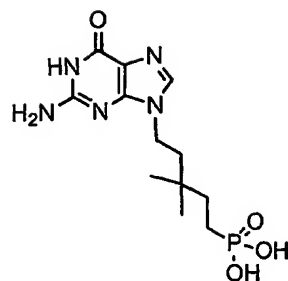
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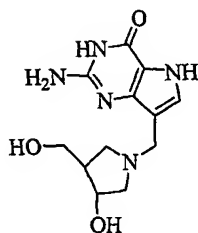
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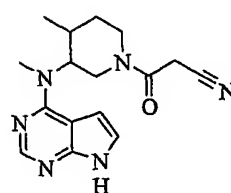
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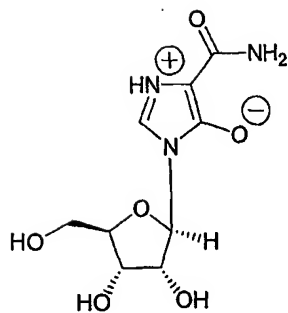


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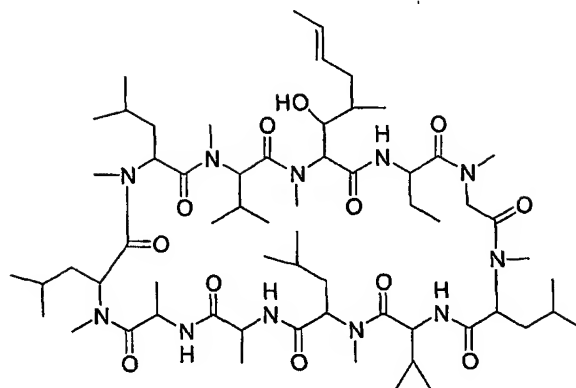


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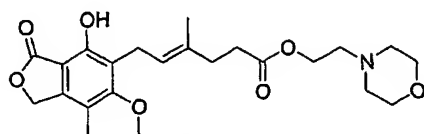
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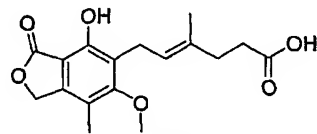
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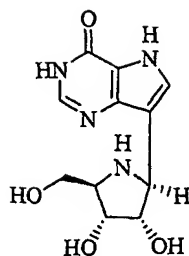


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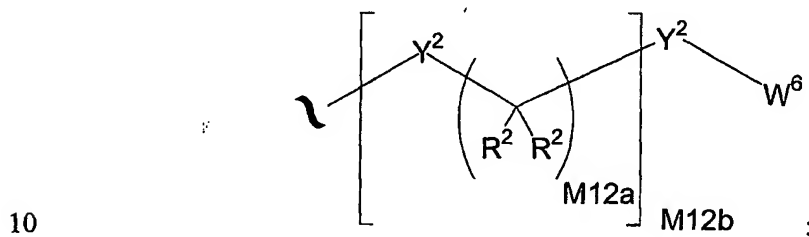


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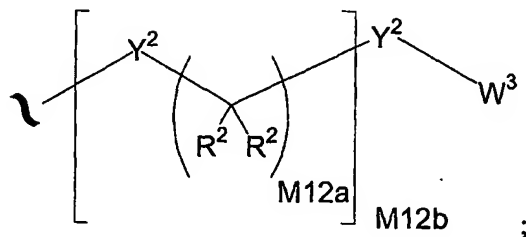
that is substituted with one or more phosphonate groups either directly or
 5 indirectly through a linker; and that is optionally substituted with one or more
 groups A^0 ; or a pharmaceutically acceptable salt thereof, wherein:

A^0 is A^1 , A^2 or W^3 with the proviso that the conjugate includes at least
 one A^1 ;

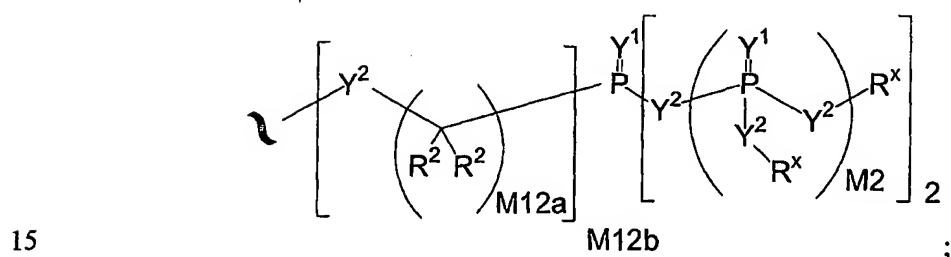
A^1 is:



A^2 is:



A^3 is:

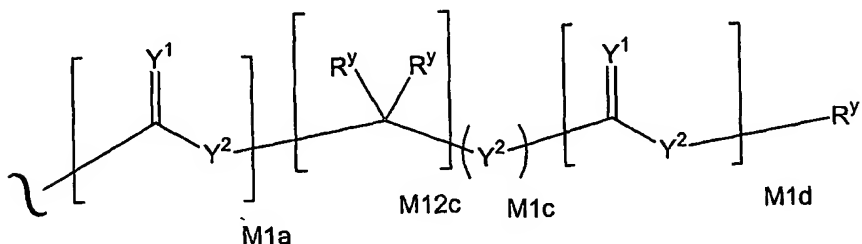


Y^1 is independently O, S, $N(R^x)$, $N(O)(R^x)$, $N(OR^x)$, $N(O)(OR^x)$, or $N(N(R^x)(R^x))$;

Y^2 is independently a bond, O, $N(R^x)$, $N(O)(R^x)$, $N(OR^x)$, $N(O)(OR^x)$, $N(N(R^x)(R^x))$, $-S(O)_{M2}-$, or $-S(O)_{M2}-S(O)_{M2}-$; and when Y^2 joins two

5 phosphorous atoms Y^2 can also be $C(R^2)(R^2)$;

R^x is independently H, R^1 , R^2 , W^3 , a protecting group, or the formula:



wherein:

R^y is independently H, W^3 , R^2 or a protecting group;

10 R^1 is independently H or alkyl of 1 to 18 carbon atoms;

R^2 is independently H, R^1 , R^3 or R^4 wherein each R^4 is independently substituted with 0 to 3 R^3 groups or taken together at a carbon atom, two R^2 groups form a ring of 3 to 8 carbons and the ring may be substituted with 0 to 3 R^3 groups;

15 R^3 is R^{3a} , R^{3b} , R^{3c} or R^{3d} , provided that when R^3 is bound to a heteroatom, then R^3 is R^{3c} or R^{3d} ;

R^{3a} is F, Cl, Br, I, $-CN$, N_3 or $-NO_2$;

R^{3b} is Y^1 ;

20 R^{3c} is $-R^x$, $-N(R^x)(R^x)$, $-SR^x$, $-S(O)R^x$, $-S(O)_2R^x$, $-S(O)(OR^x)$, $-S(O)_2(OR^x)$, $-OC(Y^1)R^x$, $-OC(Y^1)OR^x$, $-OC(Y^1)(N(R^x)(R^x))$, $-SC(Y^1)R^x$, $-SC(Y^1)OR^x$, $-SC(Y^1)(N(R^x)(R^x))$, $-N(R^x)C(Y^1)R^x$, $-N(R^x)C(Y^1)OR^x$, or $-N(R^x)C(Y^1)(N(R^x)(R^x))$;

R^{3d} is $-C(Y^1)R^x$, $-C(Y^1)OR^x$ or $-C(Y^1)(N(R^x)(R^x))$;

25 R^4 is an alkyl of 1 to 18 carbon atoms, alkenyl of 2 to 18 carbon atoms, or alkynyl of 2 to 18 carbon atoms;

R^5 is R^4 wherein each R^4 is substituted with 0 to 3 R^3 groups;

W^3 is W^4 or W^5 ;

W^4 is R^5 , $-C(Y^1)R^5$, $-C(Y^1)W^5$, $-SO_{M2}R^5$, or $-SO_{M2}W^5$;

W^5 is carbocycle or heterocycle wherein W^5 is independently substituted with 0 to 3 R^2 groups;

W^6 is W^3 independently substituted with 1, 2, or 3 A^3 groups;

M2 is 0, 1 or 2;

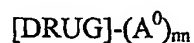
5 M12a is 1, 2, 3, 4, 5, 6, 7, 8, 9, 10, 11 or 12;

M12b is 0, 1, 2, 3, 4, 5, 6, 7, 8, 9, 10, 11 or 12;

M1a, M1c, and M1d are independently 0 or 1; and

M12c is 0, 1, 2, 3, 4, 5, 6, 7, 8, 9, 10, 11 or 12.

10 In another embodiment the invention provides a compound of the formula:



15 or a pharmaceutically acceptable salt thereof wherein;

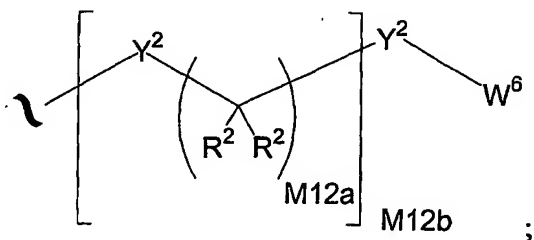
DRUG is a compound of any one of formulae 500-547;

mn is 1, 2, or 3;

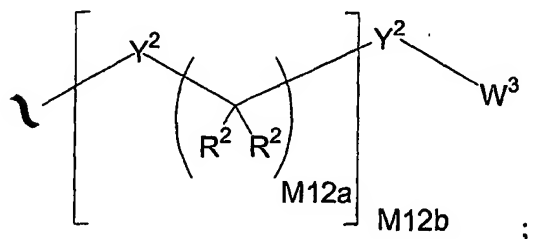
wherein:

20 A^0 is A^1 , A^2 or W^3 with the proviso that the conjugate includes at least one A^1 ;

A^1 is:

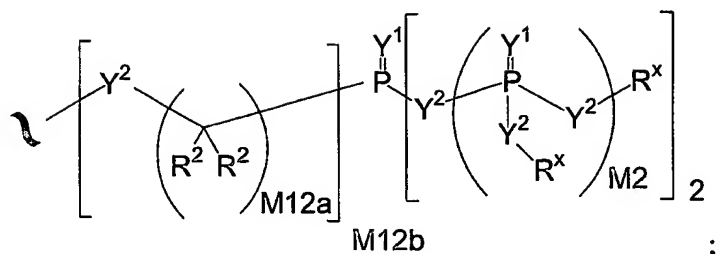


A^2 is:



25

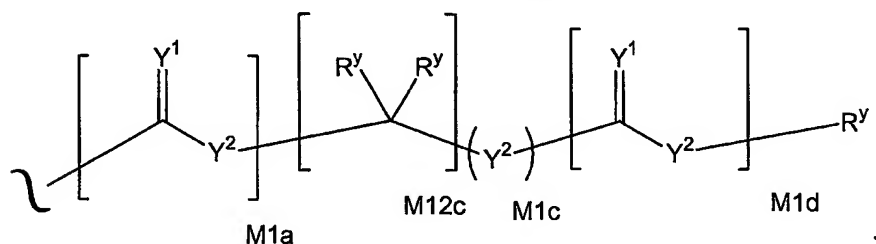
A³ is:



Y¹ is independently O, S, N(R^x), N(O)(R^x), N(OR^x), N(O)(OR^x), or N(N(R^x)(R^x));

5 Y² is independently a bond, O, N(R^x), N(O)(R^x), N(OR^x), N(O)(OR^x), N(N(R^x)(R^x)), -S(O)_{M2}-, or -S(O)_{M2}-S(O)_{M2}-; and when Y² joins two phosphorous atoms Y² can also be C(R²)(R²);

R^x is independently H, R¹, R², W³, a protecting group, or the formula:



10 wherein:

R^y is independently H, W³, R² or a protecting group;

R¹ is independently H or alkyl of 1 to 18 carbon atoms;

15 R² is independently H, R¹, R³ or R⁴ wherein each R⁴ is independently substituted with 0 to 3 R³ groups or taken together at a carbon atom, two R² groups form a ring of 3 to 8 carbons and the ring may be substituted with 0 to 3 R³ groups;

R³ is R^{3a}, R^{3b}, R^{3c} or R^{3d}, provided that when R³ is bound to a heteroatom, then R³ is R^{3c} or R^{3d};

R^{3a} is F, Cl, Br, I, -CN, N₃ or -NO₂;

20 R^{3b} is Y¹;

R^{3c} is -R^x, -N(R^x)(R^x), -SR^x, -S(O)R^x, -S(O)₂R^x, -S(O)(OR^x), -S(O)₂(OR^x), -OC(Y¹)R^x, -OC(Y¹)OR^x, -OC(Y¹)(N(R^x)(R^x)), -SC(Y¹)R^x, -SC(Y¹)OR^x, -SC(Y¹)(N(R^x)(R^x)), -N(R^x)C(Y¹)R^x, -N(R^x)C(Y¹)OR^x, or -N(R^x)C(Y¹)(N(R^x)(R^x));

R^{3d} is $-C(Y^1)R^x$, $-C(Y^1)OR^x$ or $-C(Y^1)(N(R^x)(R^x))$;

R^4 is an alkyl of 1 to 18 carbon atoms, alkenyl of 2 to 18 carbon atoms, or alkynyl of 2 to 18 carbon atoms;

R^5 is R^4 wherein each R^4 is substituted with 0 to 3 R^3 groups;

5 W^3 is W^4 or W^5 ;

W^4 is R^5 , $-C(Y^1)R^5$, $-C(Y^1)W^5$, $-SO_{M2}R^5$, or $-SO_{M2}W^5$;

W^5 is carbocycle or heterocycle wherein W^5 is independently substituted with 0 to 3 R^2 groups;

W^6 is W^3 independently substituted with 1, 2, or 3 A^3 groups;

10 $M2$ is 0, 1 or 2;

$M12a$ is 1, 2, 3, 4, 5, 6, 7, 8, 9, 10, 11 or 12;

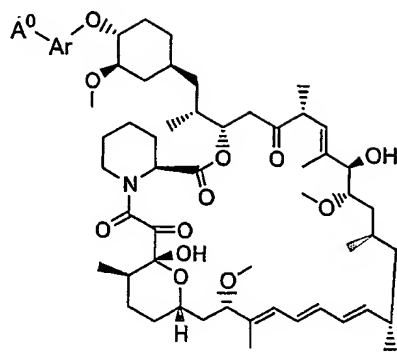
$M12b$ is 0, 1, 2, 3, 4, 5, 6, 7, 8, 9, 10, 11 or 12;

$M1a$, $M1c$, and $M1d$ are independently 0 or 1; and

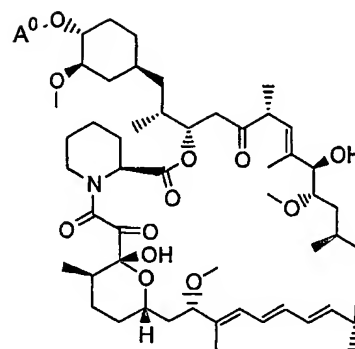
$M12c$ is 0, 1, 2, 3, 4, 5, 6, 7, 8, 9, 10, 11 or 12.

15

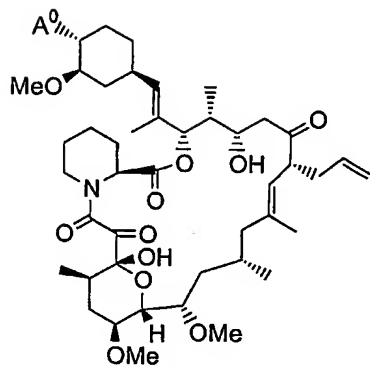
In another embodiment the invention provides a compound of any one of formulae 1-151:



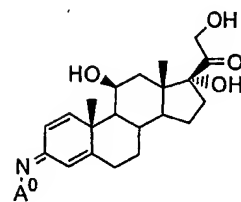
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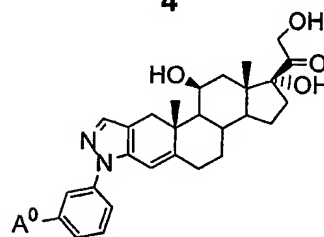
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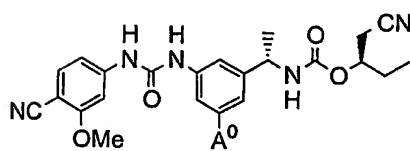
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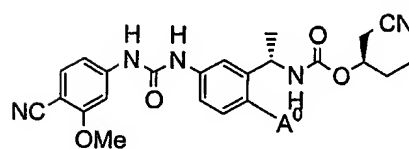
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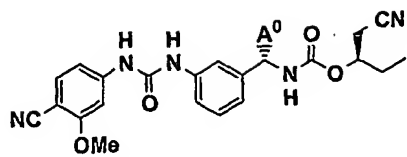
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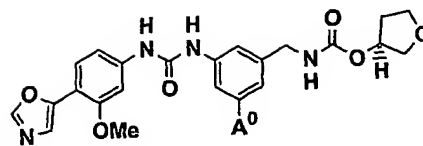
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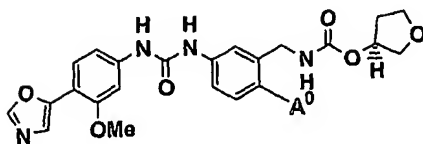
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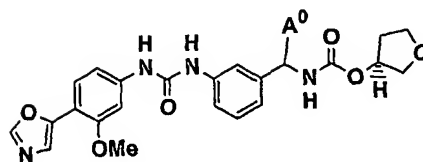
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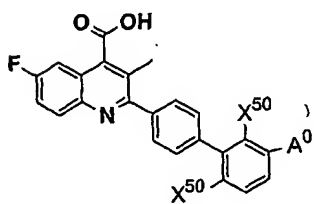
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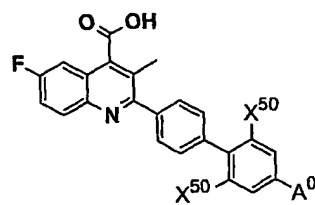
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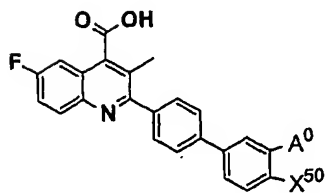
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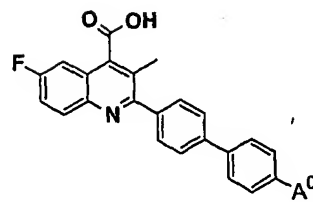
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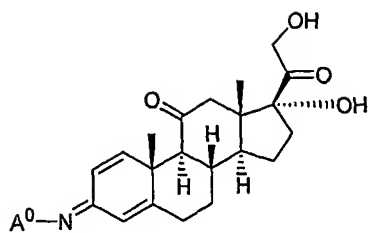
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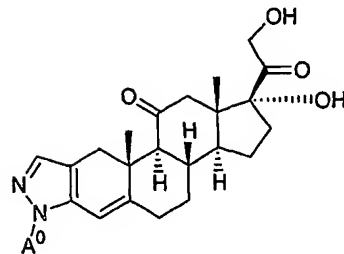
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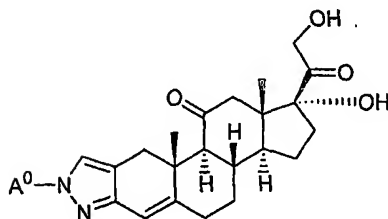
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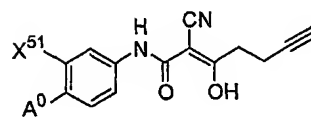
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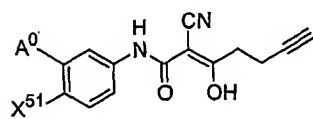
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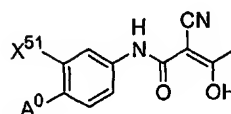
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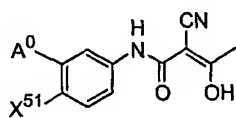
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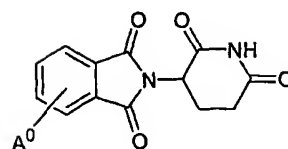
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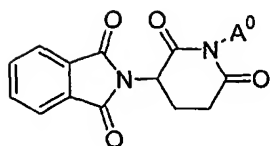
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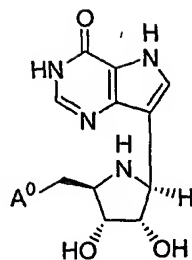
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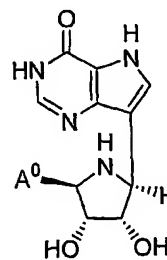
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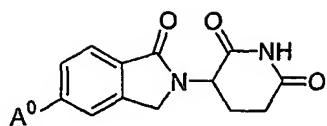
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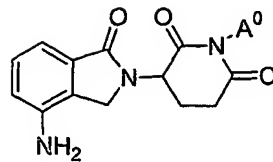
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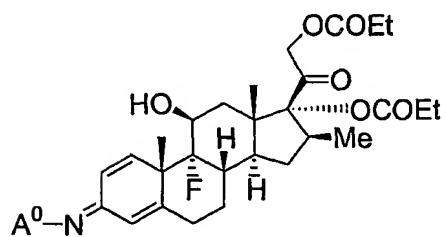


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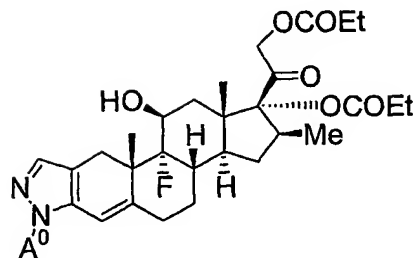


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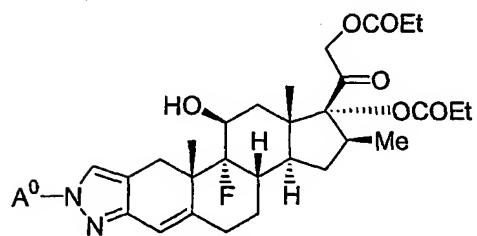
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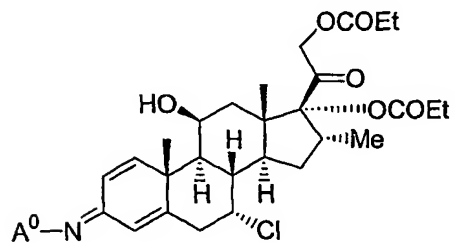
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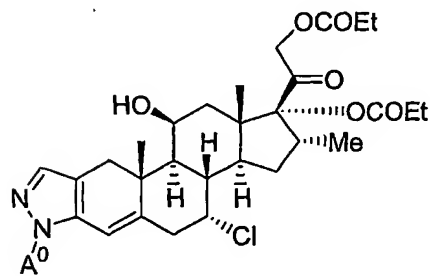


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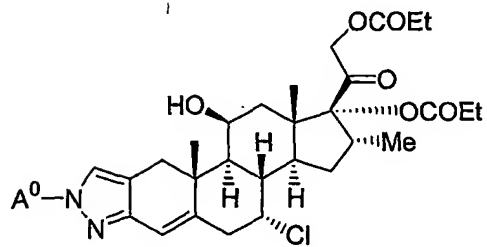


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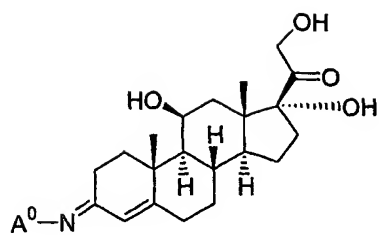
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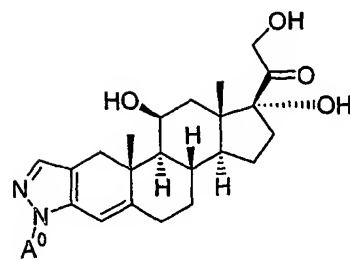
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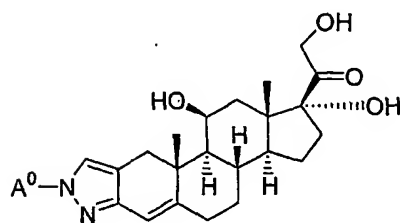
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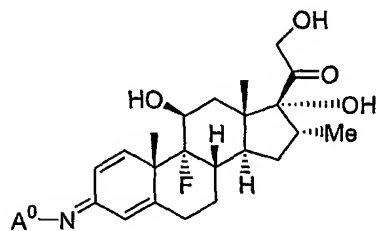
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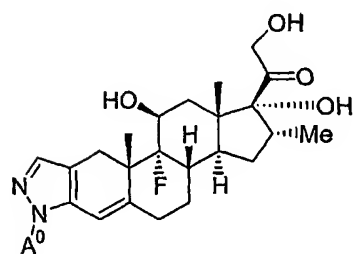


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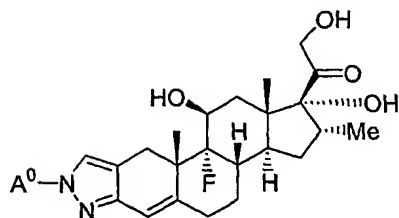


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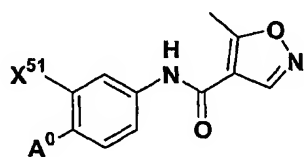
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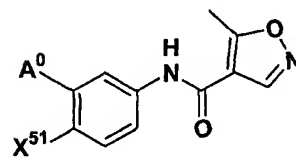
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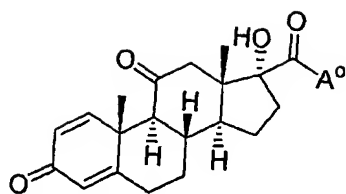


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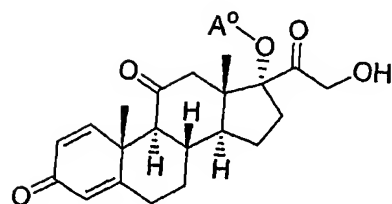


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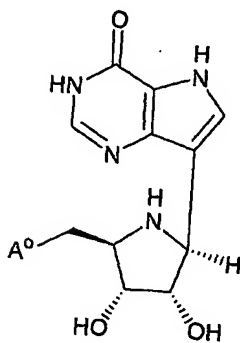
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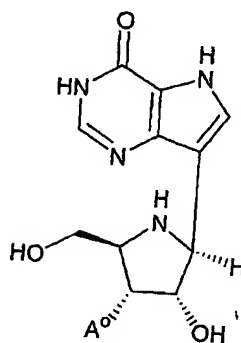
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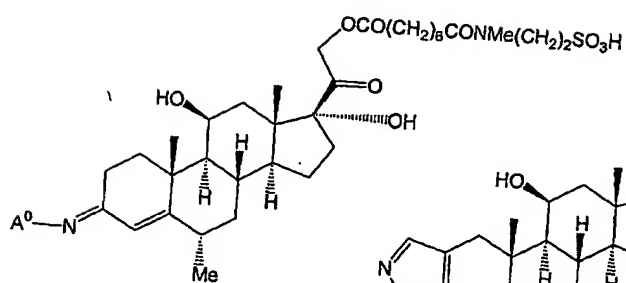


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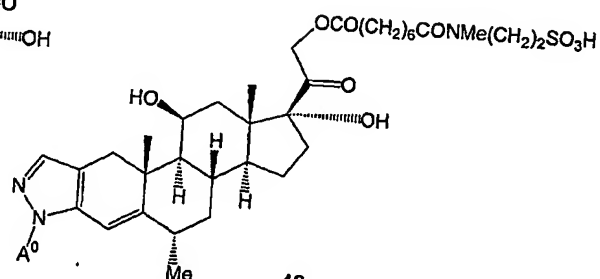


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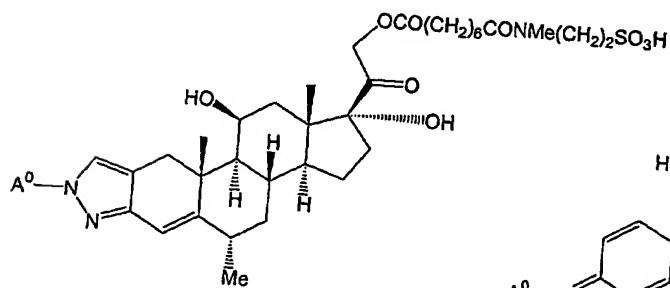
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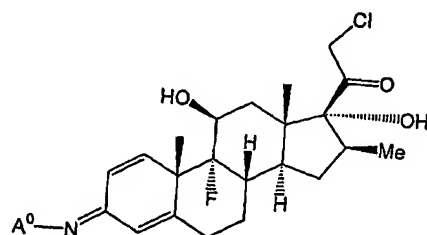
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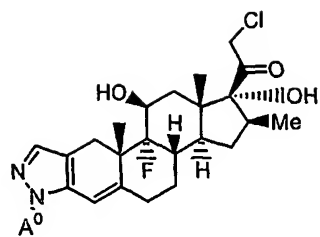


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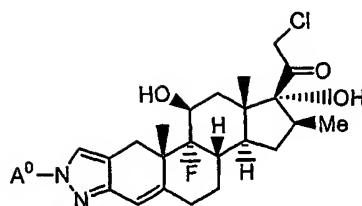


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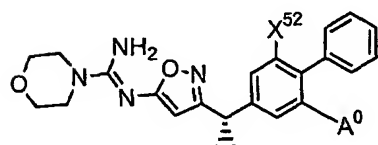
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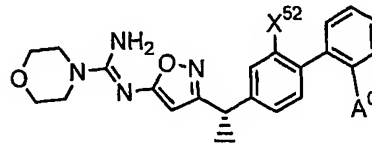
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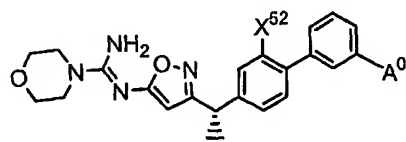
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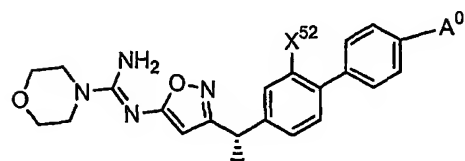
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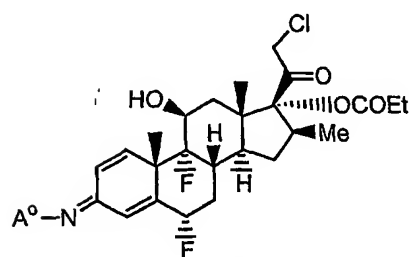


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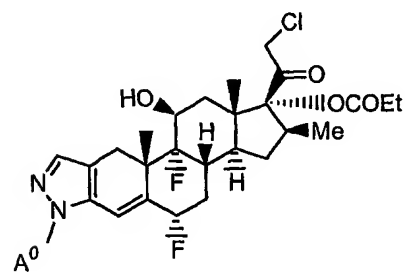


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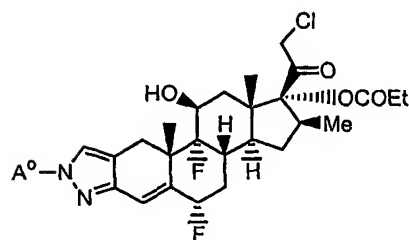


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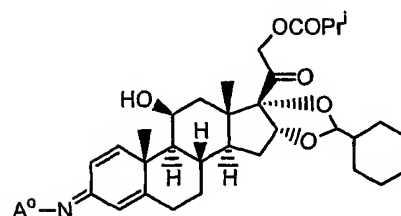


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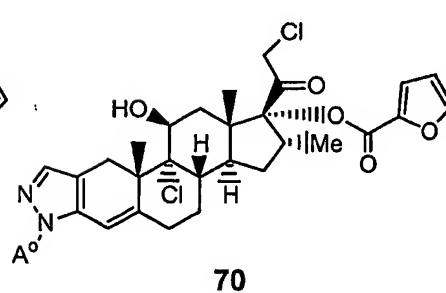
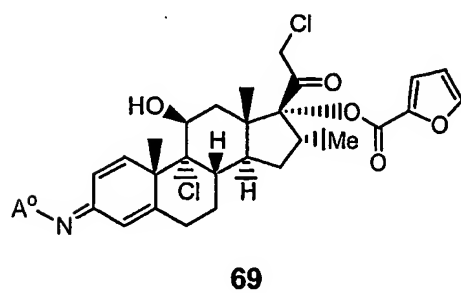
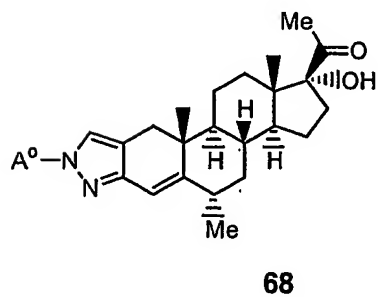
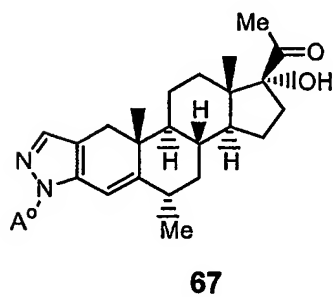
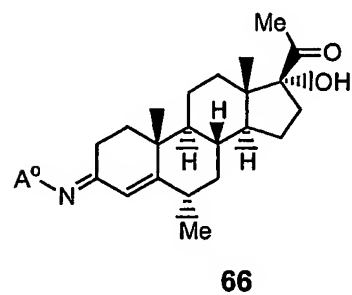
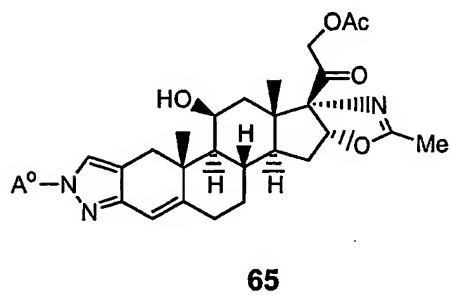
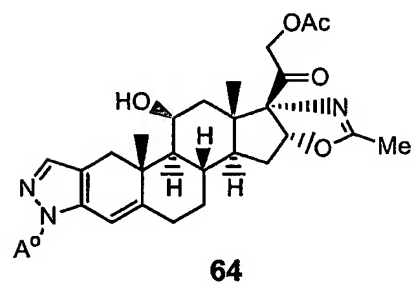
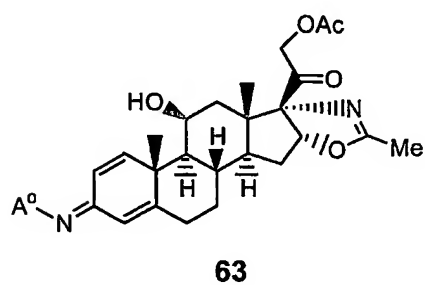
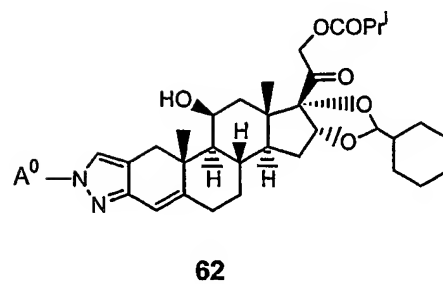
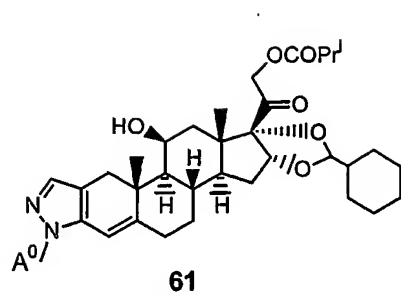
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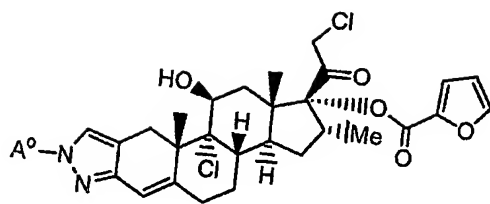


59

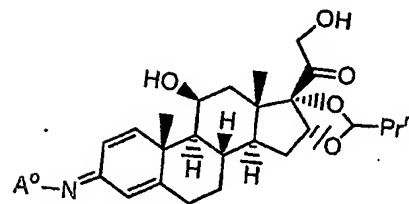


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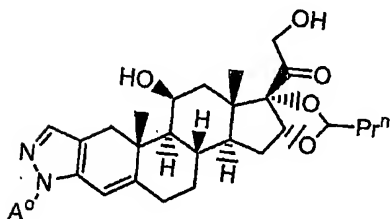




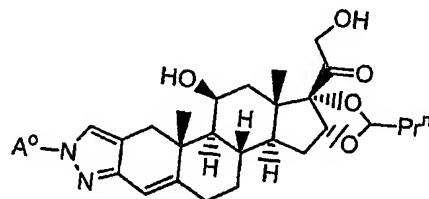
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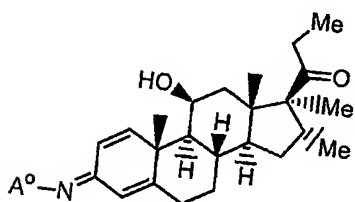
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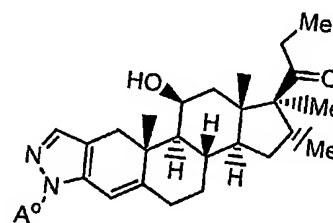
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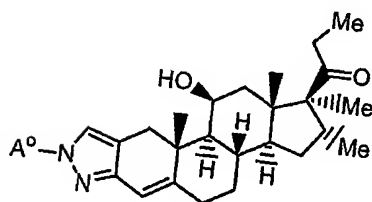
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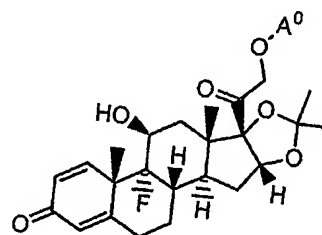
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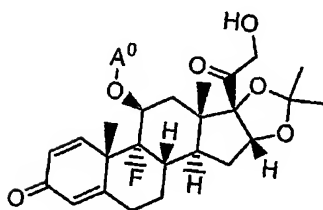
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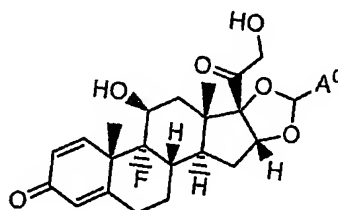
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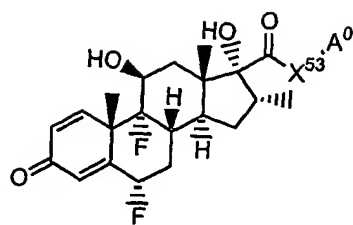
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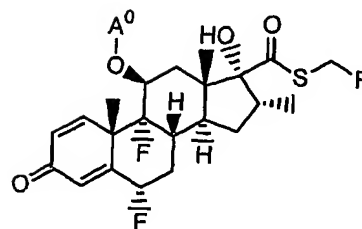
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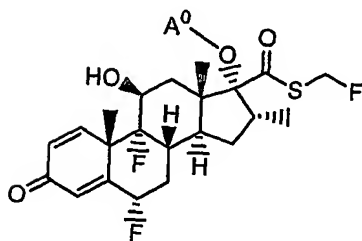
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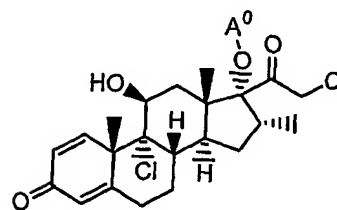
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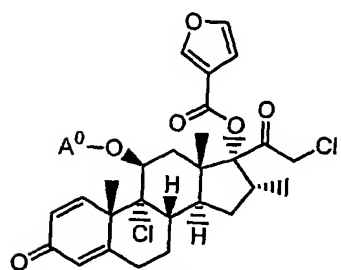
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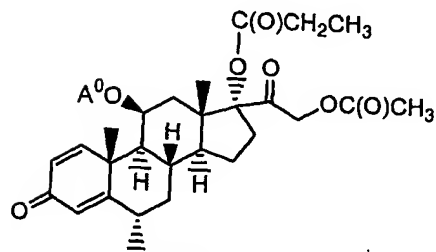
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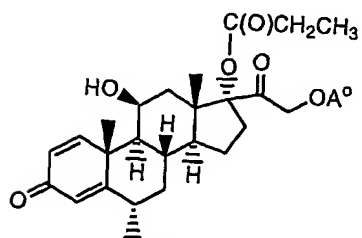
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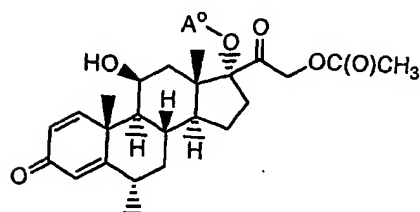
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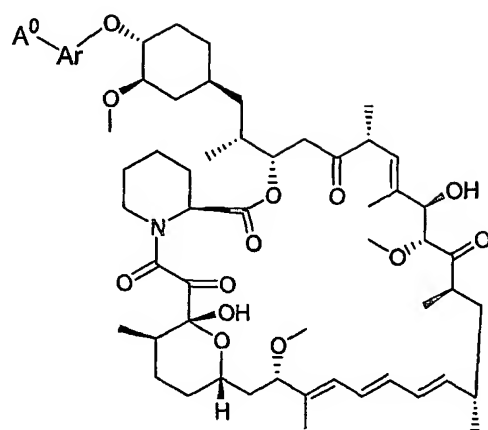
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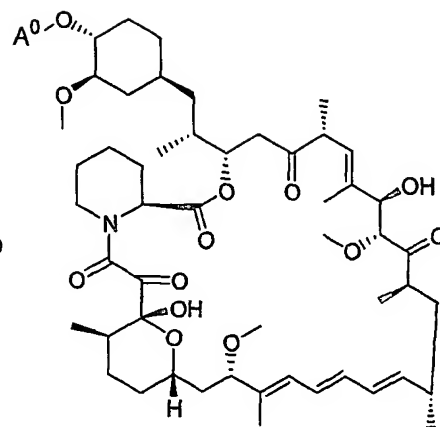
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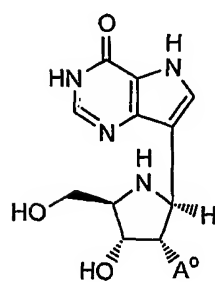
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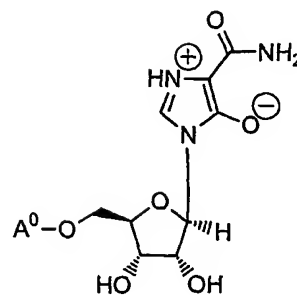
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90

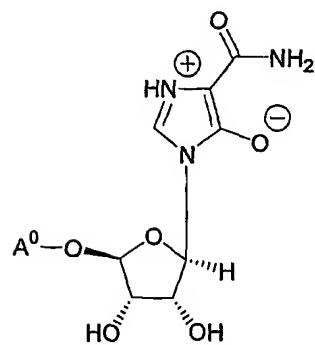


91



92

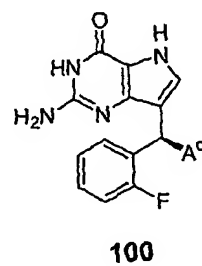
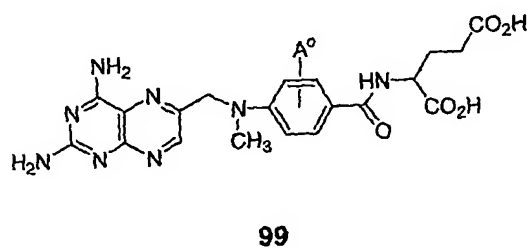
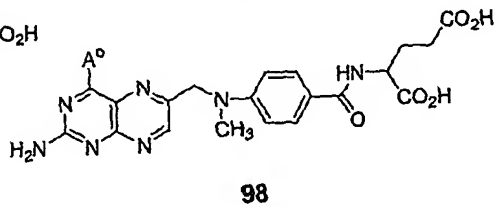
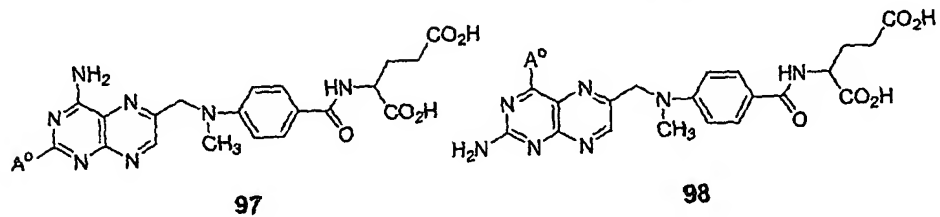
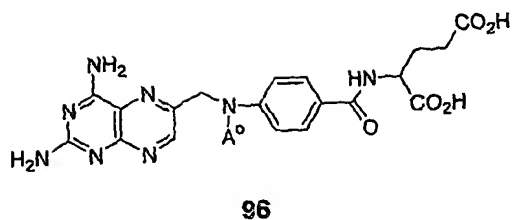
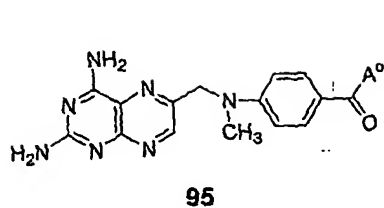
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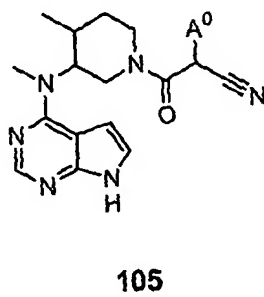
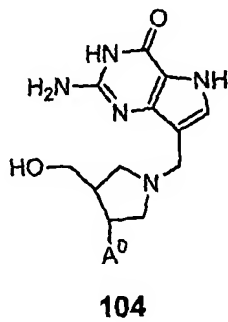
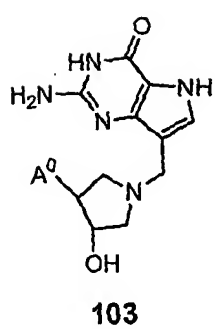
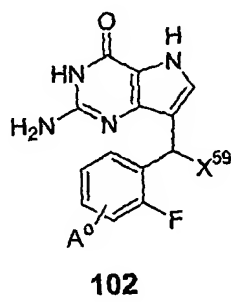
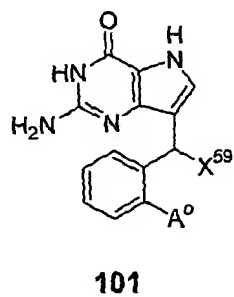
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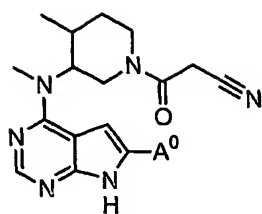
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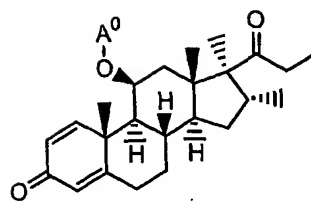
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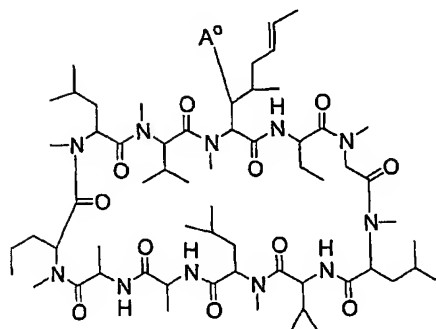
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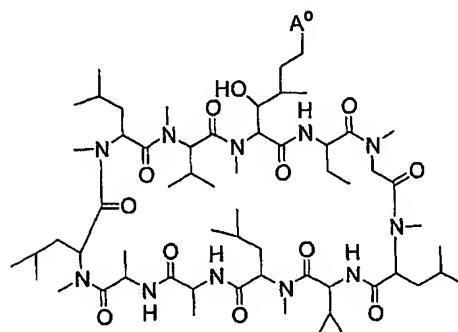
106



107

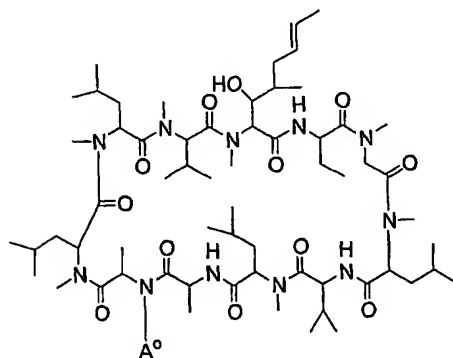


108

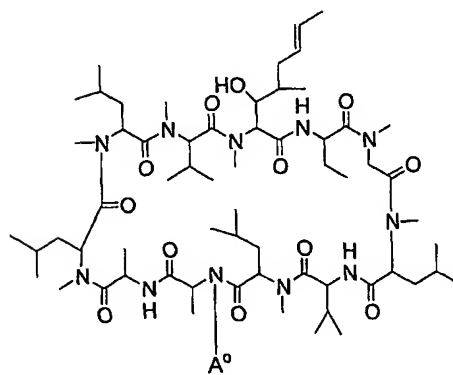


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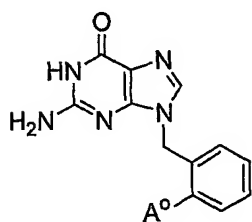
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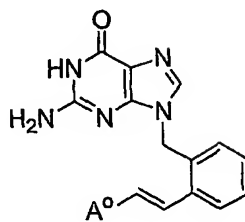
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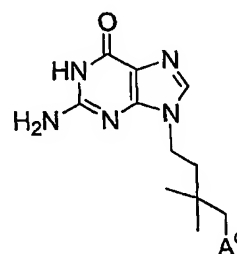
111



112

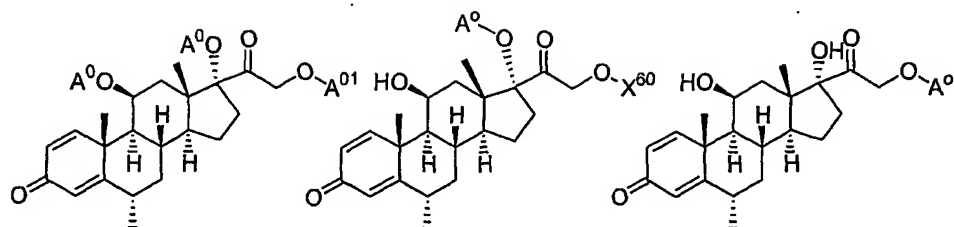


113



114

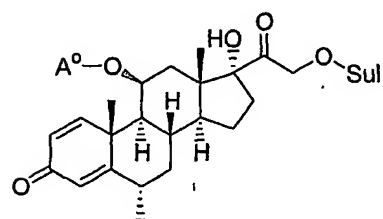
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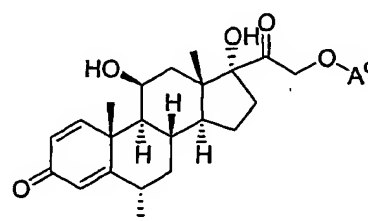
115

116

117

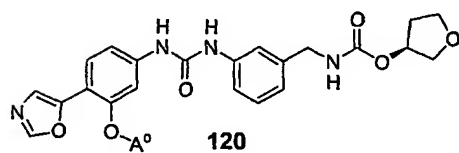
sul = CO(CH₂)₆CONMe(CH₂)₂SO₃H

118

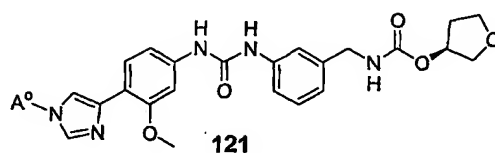


119

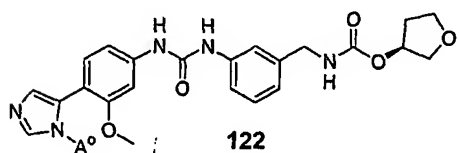
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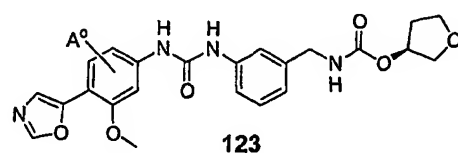
120



121

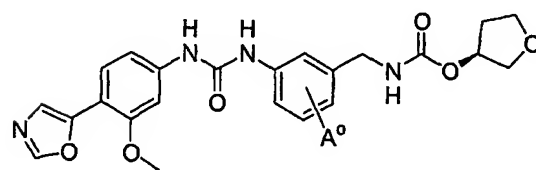


122

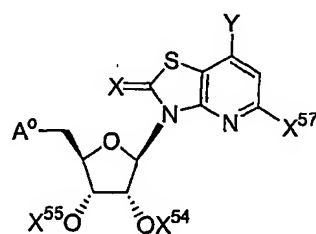


123

10

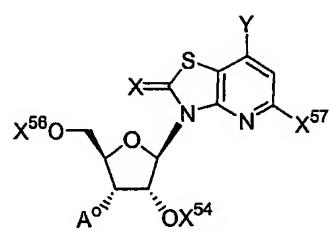


124

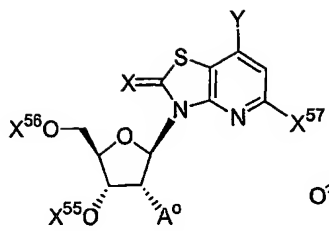


125

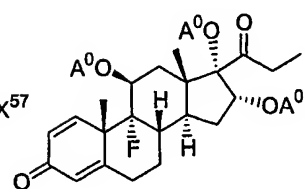
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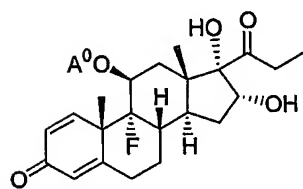
126



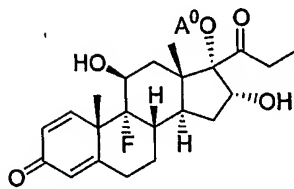
127



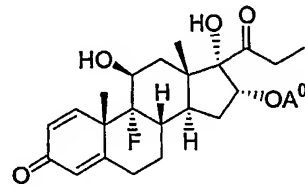
128



129

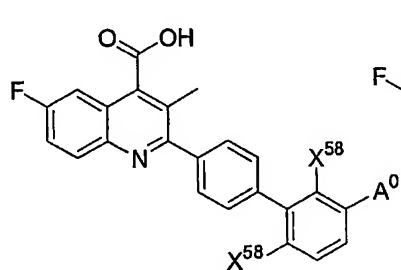


130

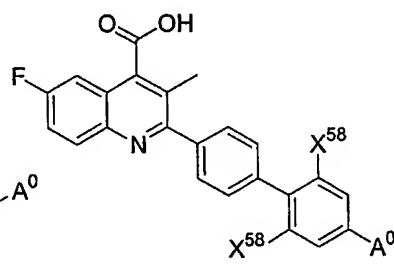


131

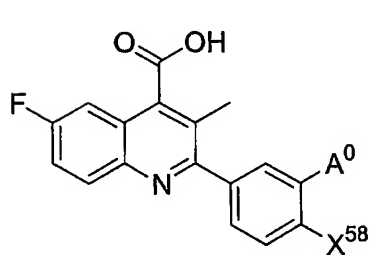
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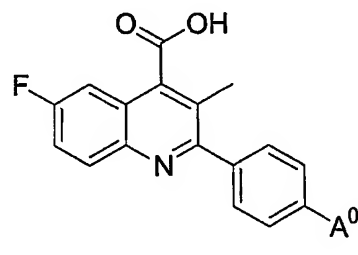
132



133

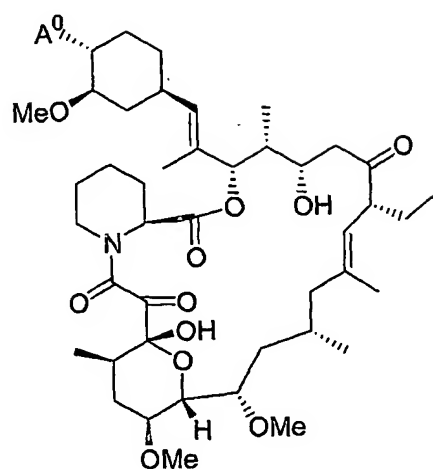
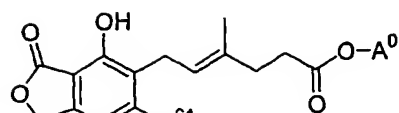
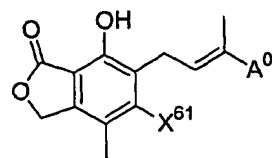
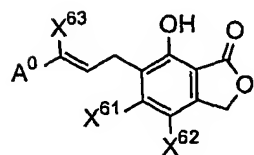
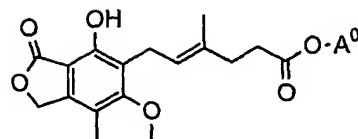


134

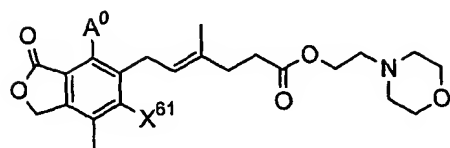
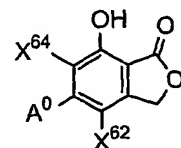
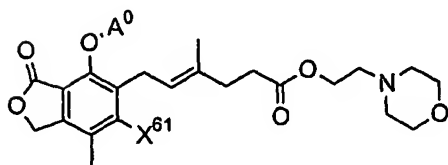
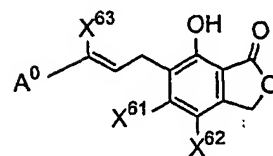


135

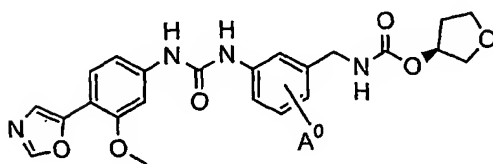
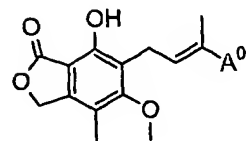
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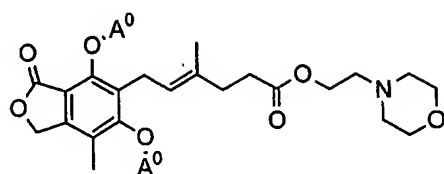
**136****137****138****139****140**

5

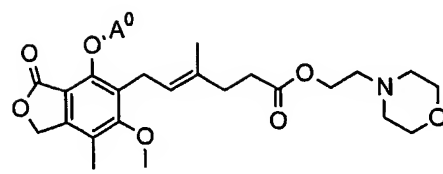
**141****142****143****144**

10

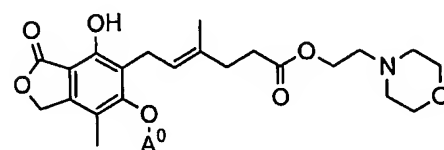
**145****146**



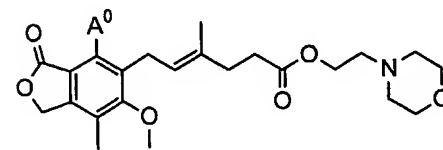
147



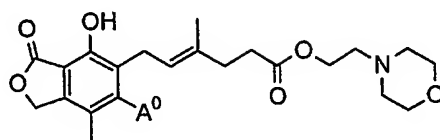
148



149



150

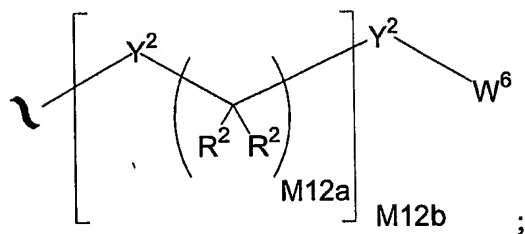


151

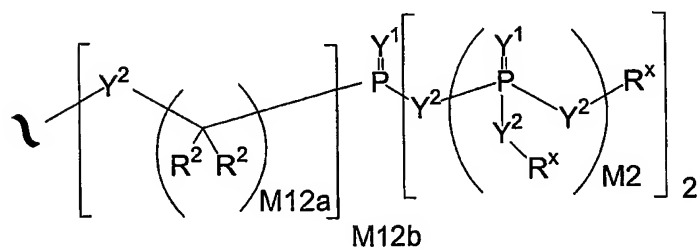
10 wherein:

A^0 is A^1 ;

A^1 is:



A^3 is:

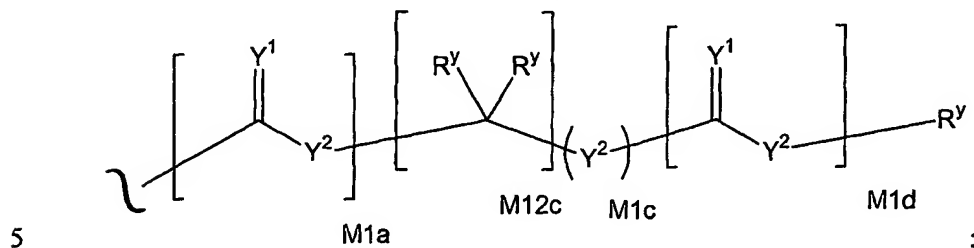


15

Y^1 is independently O, S, $N(R^x)$, $N(O)(R^x)$, $N(OR^x)$, $N(O)(OR^x)$, or $N(N(R^x)(R^x))$;

Y^2 is independently a bond, O, $N(R^x)$, $N(O)(R^x)$, $N(OR^x)$, $N(O)(OR^x)$, $N(N(R^x)(R^x))$, $-S(O)_{M2-}$, or $-S(O)_{M2-}S(O)_{M2-}$; and when Y^2 joins two phosphorous atoms Y^2 can also be $C(R^2)(R^2)$;

R^x is independently H, R^2 , W^3 , a protecting group, or the formula:



R^y is independently H, W^3 , R^2 or a protecting group;

R^1 is independently H or alkyl of 1 to 18 carbon atoms;

R^2 is independently H, R^3 or R^4 wherein each R^4 is independently substituted with 0 to 3 R^3 groups;

10 R^3 is R^{3a} , R^{3b} , R^{3c} or R^{3d} , provided that when R^3 is bound to a heteroatom, then R^3 is R^{3c} or R^{3d} ;

R^{3a} is F, Cl, Br, I, -CN, N_3 or $-NO_2$;

R^{3b} is Y^1 ;

15 R^{3c} is $-R^x$, $-N(R^x)(R^x)$, $-SR^x$, $-S(O)R^x$, $-S(O)_2R^x$, $-S(O)(OR^x)$, $-S(O)_2(OR^x)$, $-OC(Y^1)R^x$, $-OC(Y^1)OR^x$, $-OC(Y^1)(N(R^x)(R^x))$, $-SC(Y^1)R^x$, $-SC(Y^1)OR^x$, $-SC(Y^1)(N(R^x)(R^x))$, $-N(R^x)C(Y^1)R^x$, $-N(R^x)C(Y^1)OR^x$, or $-N(R^x)C(Y^1)(N(R^x)(R^x))$;

R^{3d} is $-C(Y^1)R^x$, $-C(Y^1)OR^x$ or $-C(Y^1)(N(R^x)(R^x))$;

20 R^4 is an alkyl of 1 to 18 carbon atoms, alkenyl of 2 to 18 carbon atoms, or alkynyl of 2 to 18 carbon atoms;

R^5 is R^4 wherein each R^4 is substituted with 0 to 3 R^3 groups;

R^{5a} is independently alkylene of 1 to 18 carbon atoms, alkenylene of 2 to 18 carbon atoms, or alkynylene of 2-18 carbon atoms any one of which alkylene, alkenylene or alkynylene is substituted with 0-3 R^3 groups;

25 W^3 is W^4 or W^5 ;

W^4 is R^5 , $-C(Y^1)R^5$, $-C(Y^1)W^5$, $-SO_2R^5$, or $-SO_2W^5$;

W^5 is carbocycle or heterocycle wherein W^5 is independently substituted with 0 to 3 R^2 groups;

W^6 is W^3 independently substituted with 1, 2, or 3 A^3 groups;

M2 is 0, 1 or 2;

M12a is 1, 2, 3, 4, 5, 6, 7, 8, 9, 10, 11 or 12;

M12b is 0, 1, 2, 3, 4, 5, 6, 7, 8, 9, 10, 11 or 12;

M1a, M1c, and M1d are independently 0 or 1;

5 M12c is 0, 1, 2, 3, 4, 5, 6, 7, 8, 9, 10, 11 or 12;

each X⁵⁰ is independently hydrogen, F, Cl, CF₃, CN, methyl, or *tert*-butyl;

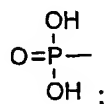
X⁵¹ is hydrogen, halo, trifluoromethyl, (C1-C3)alkyl, cyano, or (C1-C3)alkoxy;

10 X⁵² is hydrogen, fluoro, chloro, bromo, methyl, or trifluoromethyl;

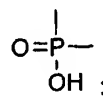
X⁵³ is -O-, or -S-;

X⁵⁴ and X⁵⁵ are independently selected from hydrogen or a C₁-C₁₈ acyl;

X⁵⁶ is hydrogen, a C₁-C₁₈ acyl, or



15 or X⁵⁴ is hydrogen and together X⁵⁵ and X⁵⁶ are



X⁵⁷ is H, amino, hydroxy, or a halogen selected from Cl and Br;

X⁵⁸ is hydrogen, F, Cl, CF₃, cyano, methyl, or *t*-butyl;

X⁵⁹ is hydrogen, CH₂OH;

20 X⁶⁰ is CO(CH₂)₆CONMe(CH₂)₂SO₃H;

X⁶² is methyl, chloro, or trifluoromethyl;

X⁶³ is H, methyl, ethyl, cyclopropyl, vinyl, or trifluoromethyl;

X⁶⁴ is H, methyl, ethyl, cyclopropyl, chloro, vinyl, allyl, 3-methyl-1-buten-1-yl;

25 X⁶⁵ is hydrogen or F; and

Ar is aryl or heteroaryl.

The invention provides a pharmaceutical composition comprising an effective amount of a compound of the invention, or a pharmaceutically acceptable salt thereof, in combination with a pharmaceutically acceptable

30 diluent or carrier.

This invention pertains to a method of increasing cellular accumulation and retention of drug compounds, thus improving their therapeutic and diagnostic value, comprising linking the compound to one or more phosphonate groups.

5 The invention also provides a method for the maintenance of immunosuppression, for example, following transplant surgery, comprising administering to an animal (e.g. a mammal) an effective amount of a compound of the invention.

10 The invention also provides a method for modulating an immune response *in vitro* or *in vivo* comprising contacting a sample in need of such treatment with a compound of the invention.

The invention also provides a method of inhibiting an immune response in an animal (e.g. a mammal), comprising administering an effective amount of a compound of the invention to the animal.

15 The invention also provides a method of treating the symptoms or effects of an autoimmune disease (e.g. psoriasis, rheumatoid arthritis, lupus erythematosus, multiple sclerosis, diabetes, Chron's disease, etc.) in an animal (e.g. a mammal), comprising administering an effective amount of a compound of the invention to the animal.

20 The invention also provides a method of treating the symptoms or effects of transplant rejection in an animal (e.g. a mammal), comprising administering an effective amount of a compound of the invention to the animal.

25 The invention also provides a method for inhibiting the proliferation of human T cells and/or downregulating the production of Th1 or Th2 type cytokines in an animal (e.g. a mammal) comprising administering a compound of the invention to the animal.

The invention also provides a method for treating atopic dermatitis in an animal (e.g. a mammal), comprising administering an effective amount of a compound of the invention to the animal.

30 The invention also provides a method for inhibiting one or more T-lymphocyte functions in an animal (e.g. a mammal), comprising administering an effective amount of a compound of the invention to the animal.

The invention also provides a method for inhibiting dihydroorotate dehydrogenase in an animal (e.g. a mammal), comprising administering an effective amount of a compound of the invention to the animal.

5 The invention also provides a compound of the invention for use in medical therapy (preferably for use in the maintenance of immunosuppression following transplant surgery, inhibiting an immune response, treating an autoimmune disease, treating atopic dermatitis, inhibiting the proliferation of human T cells, or downregulating the production of Th1 or Th2 type cytokines), as well as the use of a compound of the invention for the manufacture of a
10 medicament useful for maintenance of immunosuppression following transplant surgery in an animal (e.g. a mammal). The invention also provides the use of a compound of the invention for the manufacture of a medicament useful for inhibiting an immune response in an animal (e.g. a mammal). The invention also provides the use of a compound of the invention for the manufacture of a
15 medicament useful for treating an autoimmune disease in an animal (e.g. a mammal). The invention also provides the use of a compound of the invention for the manufacture of a medicament useful for inhibiting the proliferation of human T cells or downregulating the production of Th1 or Th2 type cytokines in an animal.

20 The invention also provides a method for the maintenance of immunosuppression, for example, following transplant surgery, comprising administering to an animal (e.g., a mammal) an effective amount of a compound of the invention.

The invention also provides processes and novel intermediates disclosed
25 herein which are useful for preparing compounds of the invention. Some of the compounds of the invention are useful to prepare other compounds of the invention.

DETAILED DESCRIPTION OF THE INVENTION

Reference will now be made in detail to certain claims of the invention,
30 examples of which are illustrated in the accompanying structures and formulas. While the invention will be described in conjunction with the enumerated claims, it will be understood that they are not intended to limit the invention to those claims. On the contrary, the invention is intended to cover all alternatives,

modifications, and equivalents, which may be included within the scope of the present invention as defined by the claims.

DEFINITIONS

Unless stated otherwise, the following terms and phrases as used herein
5 are intended to have the following meanings:

When tradenames are used herein, applicants intend to independently include the tradename product and the active pharmaceutical ingredient(s) of the tradename product.

“Bioavailability” is the degree to which the pharmaceutically active agent
10 becomes available to the target tissue after the agent's introduction into the body. Enhancement of the bioavailability of a pharmaceutically active agent can provide a more efficient and effective treatment for patients because, for a given dose, more of the pharmaceutically active agent will be available at the targeted tissue sites.

15 The terms “phosphonate” and “phosphonate group” include functional groups or moieties within a molecule that comprises a phosphorous that is 1) single-bonded to a carbon, 2) double-bonded to a heteroatom, 3) single-bonded to a heteroatom, and 4) single-bonded to another heteroatom, wherein each heteroatom can be the same or different. The terms “phosphonate” and
20 “phosphonate group” also include functional groups or moieties that comprise a phosphorous in the same oxidation state as the phosphorous described above, as well as functional groups or moieties that comprise a prodrug moiety that can separate from a compound so that the compound retains a phosphorous having the characteristics described above. For example, the terms “phosphonate” and
25 “phosphonate group” include phosphonic acid, phosphonic monoester, phosphonic diester, phosphonamidate, and phosphonthioate functional groups. In one specific embodiment of the invention, the terms “phosphonate” and “phosphonate group” include functional groups or moieties within a molecule that comprises a phosphorous that is 1) single-bonded to a carbon, 2) double-
30 bonded to an oxygen, 3) single-bonded to an oxygen, and 4) single-bonded to another oxygen, as well as functional groups or moieties that comprise a prodrug moiety that can separate from a compound so that the compound retains a phosphorous having such characteristics. In another specific embodiment of the

invention, the terms “phosphonate” and “phosphonate group” include functional groups or moieties within a molecule that comprises a phosphorous that is 1) single-bonded to a carbon, 2) double-bonded to an oxygen, 3) single-bonded to an oxygen or nitrogen, and 4) single-bonded to another oxygen or nitrogen, as well as functional groups or moieties that comprise a prodrug moiety that can separate from a compound so that the compound retains a phosphorous having such characteristics.

The term “prodrug” as used herein refers to any compound that when administered to a biological system generates the drug substance, i.e. active ingredient, as a result of spontaneous chemical reaction(s), enzyme catalyzed chemical reaction(s), photolysis, and/or metabolic chemical reaction(s). A prodrug is thus a covalently modified analog or latent form of a therapeutically-active compound.

“Prodrug moiety” refers to a labile functional group which separates from the active inhibitory compound during metabolism, systemically, inside a cell, by hydrolysis, enzymatic cleavage, or by some other process (Bundgaard, Hans, “Design and Application of Prodrugs” in A Textbook of Drug Design and Development (1991), P. Krogsgaard-Larsen and H. Bundgaard, Eds. Harwood Academic Publishers, pp. 113-191). Enzymes which are capable of an enzymatic activation mechanism with the phosphonate prodrug compounds of the invention include, but are not limited to, amidases, esterases, microbial enzymes, phospholipases, cholinesterases, and phosphotases. Prodrug moieties can serve to enhance solubility, absorption and lipophilicity to optimize drug delivery, bioavailability and efficacy. A prodrug moiety may include an active metabolite or drug itself.

Exemplary prodrug moieties include the hydrolytically sensitive or labile acyloxymethyl esters $-\text{CH}_2\text{OC}(=\text{O})\text{R}^9$ and acyloxymethyl carbonates $-\text{CH}_2\text{OC}(=\text{O})\text{OR}^9$ where R^9 is C_1 - C_6 alkyl, C_1 - C_6 substituted alkyl, C_6 - C_{20} aryl or C_6 - C_{20} substituted aryl. The acyloxyalkyl ester was first used as a prodrug strategy for carboxylic acids and then applied to phosphates and phosphonates by Farquhar et al. (1983) *J. Pharm. Sci.* 72: 324; also US Patent Nos. 4816570, 4968788, 5663159 and 5792756. Subsequently, the acyloxyalkyl ester was used to deliver phosphonic acids across cell membranes and to enhance oral

bioavailability. A close variant of the acyloxyalkyl ester, the alkoxy-carbonyloxyalkyl ester (carbonate), may also enhance oral bioavailability as a prodrug moiety in the compounds of the combinations of the invention. An exemplary acyloxymethyl ester is pivaloyloxymethoxy, (POM)

- 5 $-\text{CH}_2\text{OC}(=\text{O})\text{C}(\text{CH}_3)_3$. An exemplary acyloxymethyl carbonate prodrug moiety is pivaloyloxymethylcarbonate (POC) $-\text{CH}_2\text{OC}(=\text{O})\text{OC}(\text{CH}_3)_3$.

The phosphonate group may be a phosphonate prodrug moiety. The prodrug moiety may be sensitive to hydrolysis, such as, but not limited to a pivaloyloxymethyl carbonate (POC) or POM group. Alternatively, the prodrug
10 moiety may be sensitive to enzymatic potentiated cleavage, such as a lactate ester or a phosphoramidate-ester group.

Aryl esters of phosphorus groups, especially phenyl esters, are reported to enhance oral bioavailability (De Lombaert et al. (1994) *J. Med. Chem.* 37: 498). Phenyl esters containing a carboxylic ester ortho to the phosphate have
15 also been described (Khamnei and Torrence, (1996) *J. Med. Chem.* 39:4109-4115). Benzyl esters are reported to generate the parent phosphonic acid. In some cases, substituents at the *ortho*-or *para*-position may accelerate the hydrolysis. Benzyl analogs with an acylated phenol or an alkylated phenol may generate the phenolic compound through the action of enzymes, e.g., esterases,
20 oxidases, etc., which in turn undergoes cleavage at the benzylic C-O bond to generate the phosphoric acid and the quinone methide intermediate. Examples of this class of prodrugs are described by Mitchell et al. (1992) *J. Chem. Soc. Perkin Trans. II* 2345; Glazier WO 91/19721. Still other benzylic prodrugs have been described containing a carboxylic ester-containing group attached to the
25 benzylic methylene (Glazier WO 91/19721). Thio-containing prodrugs are reported to be useful for the intracellular delivery of phosphonate drugs. These proesters contain an ethylthio group in which the thiol group is either esterified with an acyl group or combined with another thiol group to form a disulfide. Deesterification or reduction of the disulfide generates the free thio intermediate
30 which subsequently breaks down to the phosphoric acid and episulfide (Puech et al. (1993) *Antiviral Res.*, 22: 155-174; Benzaria et al. (1996) *J. Med. Chem.* 39: 4958). Cyclic phosphonate esters have also been described as prodrugs of phosphorus-containing compounds (Erion et al., US Patent No. 6312662).

“Protecting group” refers to a moiety of a compound that masks or alters the properties of a functional group or the properties of the compound as a whole. Chemical protecting groups and strategies for protection/deprotection are well known in the art. *See e.g.*, Protective Groups in Organic Chemistry,

5 Theodora W. Greene, John Wiley & Sons, Inc., New York, 1991. Protecting groups are often utilized to mask the reactivity of certain functional groups, to assist in the efficiency of desired chemical reactions, e.g., making and breaking chemical bonds in an ordered and planned fashion. Protection of functional groups of a compound alters other physical properties besides the reactivity of
10 the protected functional group, such as the polarity, lipophilicity (hydrophobicity), and other properties which can be measured by common analytical tools. Chemically protected intermediates may themselves be biologically active or inactive.

Protected compounds may also exhibit altered, and in some cases,
15 optimized properties *in vitro* and *in vivo*, such as passage through cellular membranes and resistance to enzymatic degradation or sequestration. In this role, protected compounds with intended therapeutic effects may be referred to as prodrugs. Another function of a protecting group is to convert the parental drug into a prodrug, whereby the parental drug is released upon conversion of
20 the prodrug *in vivo*. Because active prodrugs may be absorbed more effectively than the parental drug, prodrugs may possess greater potency *in vivo* than the parental drug. Protecting groups are removed either *in vitro*, in the instance of chemical intermediates, or *in vivo*, in the case of prodrugs. With chemical intermediates, it is not particularly important that the resulting products after
25 deprotection, e.g., alcohols, be physiologically acceptable, although in general it is more desirable if the products are pharmacologically innocuous.

Any reference to any of the compounds of the invention also includes a reference to a physiologically acceptable salt thereof. Examples of physiologically acceptable salts of the compounds of the invention include salts
30 derived from an appropriate base, such as an alkali metal (for example, sodium), an alkaline earth (for example, magnesium), ammonium and NX_4^+ (wherein X is C_1 – C_4 alkyl). Physiologically acceptable salts of an hydrogen atom or an amino group include salts of organic carboxylic acids such as acetic, benzoic, lactic,

fumaric, tartaric, maleic, malonic, malic, isethionic, lactobionic and succinic acids; organic sulfonic acids, such as methanesulfonic, ethanesulfonic, benzenesulfonic and p-toluenesulfonic acids; and inorganic acids, such as hydrochloric, sulfuric, phosphoric and sulfamic acids. Physiologically

- 5 acceptable salts of a compound of an hydroxy group include the anion of said compound in combination with a suitable cation such as Na^+ and NX_4^+ (wherein X is independently selected from H or a C_1 - C_4 alkyl group).

For therapeutic use, salts of active ingredients of the compounds of the invention will be physiologically acceptable, i.e. they will be salts derived from a
 10 physiologically acceptable acid or base. However, salts of acids or bases which are not physiologically acceptable may also find use, for example, in the preparation or purification of a physiologically acceptable compound. All salts, whether or not derived from a physiologically acceptable acid or base, are within the scope of the present invention.

- 15 "Alkyl" is C_1 - C_{18} hydrocarbon containing normal, secondary, tertiary or cyclic carbon atoms. Examples are methyl (Me, $-\text{CH}_3$), ethyl (Et, $-\text{CH}_2\text{CH}_3$), 1-propyl (\underline{n} -Pr, \underline{n} -propyl, $-\text{CH}_2\text{CH}_2\text{CH}_3$), 2-propyl (\underline{i} -Pr, \underline{i} -propyl, $-\text{CH}(\text{CH}_3)_2$), 1-butyl (\underline{n} -Bu, \underline{n} -butyl, $-\text{CH}_2\text{CH}_2\text{CH}_2\text{CH}_3$), 2-methyl-1-propyl (\underline{i} -Bu, \underline{i} -butyl, $-\text{CH}_2\text{CH}(\text{CH}_3)_2$), 2-butyl (\underline{s} -Bu, \underline{s} -butyl, $-\text{CH}(\text{CH}_3)\text{CH}_2\text{CH}_3$), 2-methyl-2-propyl (\underline{t} -Bu, \underline{t} -butyl, $-\text{C}(\text{CH}_3)_3$), 1-pentyl (\underline{n} -pentyl, $-\text{CH}_2\text{CH}_2\text{CH}_2\text{CH}_2\text{CH}_3$),
 20 2-pentyl ($-\text{CH}(\text{CH}_3)\text{CH}_2\text{CH}_2\text{CH}_3$), 3-pentyl ($-\text{CH}(\text{CH}_2\text{CH}_3)_2$), 2-methyl-2-butyl ($-\text{C}(\text{CH}_3)_2\text{CH}_2\text{CH}_3$), 3-methyl-2-butyl ($-\text{CH}(\text{CH}_3)\text{CH}(\text{CH}_3)_2$), 3-methyl-1-butyl ($-\text{CH}_2\text{CH}_2\text{CH}(\text{CH}_3)_2$), 2-methyl-1-butyl ($-\text{CH}_2\text{CH}(\text{CH}_3)\text{CH}_2\text{CH}_3$), 1-hexyl ($-\text{CH}_2\text{CH}_2\text{CH}_2\text{CH}_2\text{CH}_2\text{CH}_3$), 2-hexyl ($-\text{CH}(\text{CH}_3)\text{CH}_2\text{CH}_2\text{CH}_2\text{CH}_3$), 3-hexyl ($-\text{CH}(\text{CH}_2\text{CH}_3)(\text{CH}_2\text{CH}_2\text{CH}_3)$), 2-methyl-2-pentyl ($-\text{C}(\text{CH}_3)_2\text{CH}_2\text{CH}_2\text{CH}_3$), 3-methyl-2-pentyl ($-\text{CH}(\text{CH}_3)\text{CH}(\text{CH}_3)\text{CH}_2\text{CH}_3$), 4-methyl-2-pentyl ($-\text{CH}(\text{CH}_3)\text{CH}_2\text{CH}(\text{CH}_3)_2$), 3-methyl-3-pentyl ($-\text{C}(\text{CH}_3)(\text{CH}_2\text{CH}_3)_2$), 2-methyl-3-pentyl ($-\text{CH}(\text{CH}_2\text{CH}_3)\text{CH}(\text{CH}_3)_2$), 2,3-dimethyl-2-butyl ($-\text{C}(\text{CH}_3)_2\text{CH}(\text{CH}_3)_2$), 3,3-dimethyl-2-butyl ($-\text{CH}(\text{CH}_3)\text{C}(\text{CH}_3)_3$).
- 25
 30

"Alkenyl" is C_2 - C_{18} hydrocarbon containing normal, secondary, tertiary or cyclic carbon atoms with at least one site of unsaturation, i.e. a carbon-carbon,

sp^2 double bond. Examples include, but are not limited to, ethylene or vinyl ($-\text{CH}=\text{CH}_2$), allyl ($-\text{CH}_2\text{CH}=\text{CH}_2$), cyclopentenyl ($-\text{C}_5\text{H}_7$), and 5-hexenyl ($-\text{CH}_2\text{CH}_2\text{CH}_2\text{CH}_2\text{CH}=\text{CH}_2$).

“Alkynyl” is C_2 - C_{18} hydrocarbon containing normal, secondary, tertiary or cyclic carbon atoms with at least one site of unsaturation, i.e. a carbon-carbon, sp triple bond. Examples include, but are not limited to, acetylenic ($-\text{C}\equiv\text{CH}$) and propargyl ($-\text{CH}_2\text{C}\equiv\text{CH}$),

“Alkylene” refers to a saturated, branched or straight chain or cyclic hydrocarbon radical of 1-18 carbon atoms, and having two monovalent radical centers derived by the removal of two hydrogen atoms from the same or two different carbon atoms of a parent alkane. Typical alkylene radicals include, but are not limited to, methylene ($-\text{CH}_2-$), 1,2-ethyl ($-\text{CH}_2\text{CH}_2-$), 1,3-propyl ($-\text{CH}_2\text{CH}_2\text{CH}_2-$), 1,4-butyl ($-\text{CH}_2\text{CH}_2\text{CH}_2\text{CH}_2-$), and the like.

“Alkenylene” refers to an unsaturated, branched or straight chain or cyclic hydrocarbon radical of 2-18 carbon atoms, and having two monovalent radical centers derived by the removal of two hydrogen atoms from the same or two different carbon atoms of a parent alkene. Typical alkenylene radicals include, but are not limited to, 1,2-ethylene ($-\text{CH}=\text{CH}-$).

“Alkynylene” refers to an unsaturated, branched or straight chain or cyclic hydrocarbon radical of 2-18 carbon atoms, and having two monovalent radical centers derived by the removal of two hydrogen atoms from the same or two different carbon atoms of a parent alkyne. Typical alkynylene radicals include, but are not limited to, acetylene ($-\text{C}\equiv\text{C}-$), propargyl ($-\text{CH}_2\text{C}\equiv\text{C}-$), and 4-pentynyl ($-\text{CH}_2\text{CH}_2\text{CH}_2\text{C}\equiv\text{CH}-$).

“Aryl” means a monovalent aromatic hydrocarbon radical of 6-20 carbon atoms derived by the removal of one hydrogen atom from a single carbon atom of a parent aromatic ring system. Typical aryl groups include, but are not limited to, radicals derived from benzene, substituted benzene, naphthalene, anthracene, biphenyl, and the like.

“Arylalkyl” refers to an acyclic alkyl radical in which one of the hydrogen atoms bonded to a carbon atom, typically a terminal or sp^3 carbon atom, is replaced with an aryl radical. Typical arylalkyl groups include, but are not limited to, benzyl, 2-phenylethan-1-yl, naphthylmethyl, 2-naphthylethan-1-

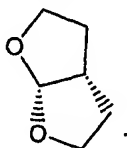
yl, naphthobenzyl, 2-naphthophenylethan-1-yl and the like. The arylalkyl group comprises 6 to 20 carbon atoms, e.g., the alkyl moiety, including alkanyl, alkenyl or alkynyl groups, of the arylalkyl group is 1 to 6 carbon atoms and the aryl moiety is 5 to 14 carbon atoms.

- 5 “Substituted alkyl”, “substituted aryl”, and “substituted arylalkyl” mean alkyl, aryl, and arylalkyl respectively, in which one or more hydrogen atoms are each independently replaced with a non-hydrogen substituent. Typical substituents include, but are not limited to, -X, -R, -O⁻, -OR, -SR, -S⁻, -NR₂, -NR₃, =NR, -CX₃, -CN, -OCN, -SCN, -N=C=O, -NCS, -NO, -NO₂, =N₂, -N₃,
 10 NC(=O)R, -C(=O)R, -C(=O)NRR, -S(=O)₂O⁻, -S(=O)₂OH, -S(=O)₂R, -OS(=O)₂OR, -S(=O)₂NR, -S(=O)R, -OP(=O)O₂RR, -P(=O)O₂RR, -P(=O)(O⁻)₂, -P(=O)(OH)₂, -C(=O)R, -C(=O)X, -C(S)R, -C(O)OR, -C(O)O⁻, -C(S)OR, -C(O)SR, -C(S)SR, -C(O)NRR, -C(S)NRR, -C(NR)NRR, where each X is independently a halogen: F, Cl, Br, or I; and each R is independently -H, alkyl,
 15 aryl, heterocycle, protecting group or prodrug moiety. Alkylene, alkenylene, and alkynylene groups may also be similarly substituted.

- “Heterocycle” as used herein includes by way of example and not limitation these heterocycles described in Paquette, Leo A.; Principles of Modern Heterocyclic Chemistry (W.A. Benjamin, New York, 1968), particularly
 20 Chapters 1, 3, 4, 6, 7, and 9; The Chemistry of Heterocyclic Compounds, A Series of Monographs (John Wiley & Sons, New York, 1950 to present), in particular Volumes 13, 14, 16, 19, and 28; and *J. Am. Chem. Soc.* (1960) 82:5566. In one specific embodiment of the invention “heterocycle” includes a “carbocycle” as defined herein, wherein one or more (e.g. 1, 2, 3, or 4) carbon
 25 atoms have been replaced with a heteroatom (e.g. O, N, or S).

- Examples of heterocycles include by way of example and not limitation pyridyl, dihydropyridyl, tetrahydropyridyl (piperidyl), thiazolyl, tetrahydrothiophenyl, sulfur oxidized tetrahydrothiophenyl, pyrimidinyl, furanyl, thienyl, pyrrolyl, pyrazolyl, imidazolyl, tetrazolyl, benzofuranyl,
 30 thianaphthalenyl, indolyl, indolenyl, quinolinyl, isoquinolinyl, benzimidazolyl, piperidinyl, 4-piperidonyl, pyrrolidinyl, 2-pyrrolidonyl, pyrrolinyl, tetrahydrofuranyl, tetrahydroquinolinyl, tetrahydroisoquinolinyl, decahydroquinolinyl, octahydroisoquinolinyl, azocinyl, triazinyl, 6H-1,2,5-

thiadiazinyl, 2H,6H-1,5,2-dithiazinyl, thienyl, thianthrenyl, pyranyl,
 isobenzofuranyl, chromenyl, xanthenyl, phenoxathinyl, 2H-pyrrolyl,
 isothiazolyl, isoxazolyl, pyrazinyl, pyridazinyl, indolizinyl, isoindolyl, 3H-
 indolyl, 1H-indazolyl, purinyl, 4H-quinolizinyl, phthalazinyl, naphthyridinyl,
 5 quinoxalinyl, quinazolinyl, cinnolinyl, pteridinyl, 4aH-carbazolyl, carbazolyl, β -
 carbolinyl, phenanthridinyl, acridinyl, pyrimidinyl, phenanthrolinyl, phenazinyl,
 phenothiazinyl, furazanyl, phenoxazinyl, isochromanyl, chromanyl,
 imidazolidinyl, imidazolyl, pyrazolidinyl, pyrazolyl, piperazinyl, indolyl,
 isoindolyl, quinuclidinyl, morpholinyl, oxazolidinyl, benzotriazolyl,
 10 benzisoxazolyl, oxindolyl, benzoxazolyl, isatinoyl, and bis-tetrahydrofuranyl:



By way of example and not limitation, carbon bonded heterocycles are
 bonded at position 2, 3, 4, 5, or 6 of a pyridine, position 3, 4, 5, or 6 of a
 15 pyridazine, position 2, 4, 5, or 6 of a pyrimidine, position 2, 3, 5, or 6 of a
 pyrazine, position 2, 3, 4, or 5 of a furan, tetrahydrofuran, thiofuran, thiophene,
 pyrrole or tetrahydropyrrole, position 2, 4, or 5 of an oxazole, imidazole or
 thiazole, position 3, 4, or 5 of an isoxazole, pyrazole, or isothiazole, position 2 or
 3 of an aziridine, position 2, 3, or 4 of an azetidine, position 2, 3, 4, 5, 6, 7, or 8
 20 of a quinoline or position 1, 3, 4, 5, 6, 7, or 8 of an isoquinoline. Still more
 typically, carbon bonded heterocycles include 2-pyridyl, 3-pyridyl, 4-pyridyl, 5-
 pyridyl, 6-pyridyl, 3-pyridazinyl, 4-pyridazinyl, 5-pyridazinyl, 6-pyridazinyl, 2-
 pyrimidinyl, 4-pyrimidinyl, 5-pyrimidinyl, 6-pyrimidinyl, 2-pyrazinyl, 3-
 pyrazinyl, 5-pyrazinyl, 6-pyrazinyl, 2-thiazolyl, 4-thiazolyl, or 5-thiazolyl.

25 By way of example and not limitation, nitrogen bonded heterocycles are
 bonded at position 1 of an aziridine, azetidine, pyrrole, pyrrolidine, 2-pyrroline,
 3-pyrroline, imidazole, imidazolidine, 2-imidazoline, 3-imidazoline, pyrazole,
 pyrazoline, 2-pyrazoline, 3-pyrazoline, piperidine, piperazine, indole, indoline,
 1H-indazole, position 2 of a isoindole, or isoindoline, position 4 of a morpholine,
 30 and position 9 of a carbazole, or β -carboline. Still more typically, nitrogen
 bonded heterocycles include 1-aziridyl, 1-azetetyl, 1-pyrrolyl, 1-imidazolyl, 1-

pyrazolyl, and 1-piperidinyl.

“Carbocycle” refers to a saturated, unsaturated or aromatic ring having 3 to 7 carbon atoms as a monocycle, 7 to 12 carbon atoms as a bicycle, and up to about 20 carbon atoms as a polycycle. Monocyclic carbocycles have 3 to 6 ring atoms, still more typically 5 or 6 ring atoms. Bicyclic carbocycles have 7 to 12 ring atoms, e.g., arranged as a bicyclo [4,5], [5,5], [5,6] or [6,6] system, or 9 or 10 ring atoms arranged as a bicyclo [5,6] or [6,6] system. Examples of monocyclic carbocycles include cyclopropyl, cyclobutyl, cyclopentyl, 1-cyclopent-1-enyl, 1-cyclopent-2-enyl, 1-cyclopent-3-enyl, cyclohexyl, 1-cyclohex-1-enyl, 1-cyclohex-2-enyl, 1-cyclohex-3-enyl, phenyl, spiryl and naphthyl.

“Linker” or “link” refers to a chemical moiety comprising a covalent bond or a chain or group of atoms that covalently attaches a phosphonate group to a drug. Linkers include portions of substituents A¹ and A³, which include moieties such as: repeating units of alkyloxy (e.g., polyethylenoxy, PEG, polymethyleneoxy) and alkylamino (e.g., polyethyleneamino, Jeffamine™); and diacid ester and amides including succinate, succinamide, diglycolate, malonate, and caproamide.

The term “chiral” refers to molecules which have the property of non-superimposability of the mirror image partner, while the term “achiral” refers to molecules which are superimposable on their mirror image partner.

The term “stereoisomers” refers to compounds which have identical chemical constitution, but differ with regard to the arrangement of the atoms or groups in space.

“Diastereomer” refers to a stereoisomer with two or more centers of chirality and whose molecules are not mirror images of one another. Diastereomers have different physical properties, e.g., melting points, boiling points, spectral properties, and reactivities. Mixtures of diastereomers may separate under high resolution analytical procedures such as electrophoresis and chromatography.

“Enantiomers” refer to two stereoisomers of a compound which are non-superimposable mirror images of one another.

The term "treatment" or "treating," to the extent it relates to a disease or condition includes preventing the disease or condition from occurring, inhibiting the disease or condition, eliminating the disease or condition, and/or relieving one or more symptoms of the disease or condition.

5 Stereochemical definitions and conventions used herein generally follow S. P. Parker, Ed., McGraw-Hill Dictionary of Chemical Terms (1984) McGraw-Hill Book Company, New York; and Eliel, E. and Wilen, S., Stereochemistry of Organic Compounds (1994) John Wiley & Sons, Inc., New York. Many organic compounds exist in optically active forms, i.e., they have the ability to rotate the
10 plane of plane-polarized light. In describing an optically active compound, the prefixes D and L or R and S are used to denote the absolute configuration of the molecule about its chiral center(s). The prefixes d and l or (+) and (-) are employed to designate the sign of rotation of plane-polarized light by the compound, with (-) or l meaning that the compound is levorotatory. A
15 compound prefixed with (+) or d is dextrorotatory. For a given chemical structure, these stereoisomers are identical except that they are mirror images of one another. A specific stereoisomer may also be referred to as an enantiomer, and a mixture of such isomers is often called an enantiomeric mixture. A 50:50 mixture of enantiomers is referred to as a racemic mixture or a racemate, which
20 may occur where there has been no stereoselection or stereospecificity in a chemical reaction or process. The terms "racemic mixture" and "racemate" refer to an equimolar mixture of two enantiomeric species, devoid of optical activity.

Protecting Groups

In the context of the present invention, protecting groups include prodrug
25 moieties and chemical protecting groups.

Protecting groups are available, commonly known and used, and are optionally used to prevent side reactions with the protected group during synthetic procedures, i.e. routes or methods to prepare the compounds of the invention. For the most part the decision as to which groups to protect, when to
30 do so, and the nature of the chemical protecting group "PG" will be dependent upon the chemistry of the reaction to be protected against (e.g., acidic, basic, oxidative, reductive or other conditions) and the intended direction of the synthesis. The PG groups do not need to be, and generally are not, the same if

the compound is substituted with multiple PG. In general, PG will be used to protect functional groups such as carboxyl, hydroxyl, thio, or amino groups and to thus prevent side reactions or to otherwise facilitate the synthetic efficiency. The order of deprotection to yield free, deprotected groups is dependent upon the intended direction of the synthesis and the reaction conditions to be encountered, and may occur in any order as determined by the artisan.

Various functional groups of the compounds of the invention may be protected. For example, protecting groups for -OH groups (whether hydroxyl, carboxylic acid, phosphonic acid, or other functions) include "ether- or ester-forming groups". Ether- or ester-forming groups are capable of functioning as chemical protecting groups in the synthetic schemes set forth herein. However, some hydroxyl and thio protecting groups are neither ether- nor ester-forming groups, as will be understood by those skilled in the art, and are included with amides, discussed below.

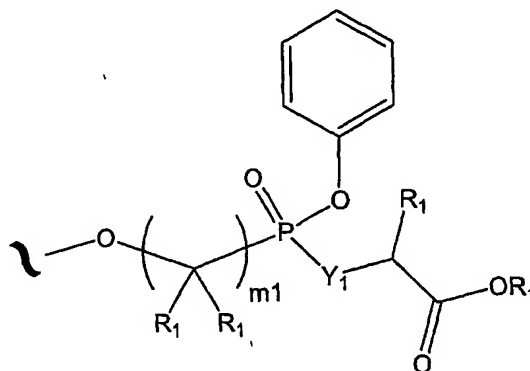
A very large number of hydroxyl protecting groups and amide-forming groups and corresponding chemical cleavage reactions are described in Protective Groups in Organic Synthesis, Theodora W. Greene (John Wiley & Sons, Inc., New York, 1991, ISBN 0-471-62301-6) ("Greene"). See also Kocienski, Philip J.; Protecting Groups (Georg Thieme Verlag Stuttgart, New York, 1994), which is incorporated by reference in its entirety herein. In particular Chapter 1, Protecting Groups: An Overview, pages 1-20, Chapter 2, Hydroxyl Protecting Groups, pages 21-94, Chapter 3, Diol Protecting Groups, pages 95-117, Chapter 4, Carboxyl Protecting Groups, pages 118-154, Chapter 5, Carbonyl Protecting Groups, pages 155-184. For protecting groups for carboxylic acid, phosphonic acid, phosphonate, sulfonic acid and other protecting groups for acids see Greene as set forth below. Such groups include by way of example and not limitation, esters, amides, hydrazides, and the like.

Ether- and Ester-forming protecting groups

Ester-forming groups include: (1) phosphonate ester-forming groups, such as phosphoramidate esters, phosphorothioate esters, phosphonate esters, and phosphon-bis-amidates; (2) carboxyl ester-forming groups, and (3) sulphur ester-forming groups, such as sulphonate, sulfate, and sulfinates.

The phosphonate moieties of the compounds of the invention may or may not be prodrug moieties, i.e. they may or may not be susceptible to hydrolytic or enzymatic cleavage or modification. Certain phosphonate moieties are stable under most or nearly all metabolic conditions. For example, a dialkylphosphonate, where the alkyl groups are two or more carbons, may have appreciable stability *in vivo* due to a slow rate of hydrolysis.

Within the context of phosphonate prodrug moieties, a large number of structurally-diverse prodrugs have been described for phosphonic acids (Freeman and Ross in Progress in Medicinal Chemistry 34: 112-147 (1997)) and are included within the scope of the present invention. An exemplary phosphonate ester-forming group is the phenyl carbocycle in substructure A₃ having the formula:



wherein R₁ may be H or C₁-C₁₂ alkyl; m₁ is 1, 2, 3, 4, 5, 6, 7 or 8, and the phenyl carbocycle is substituted with 0 to 3 R₂ groups. Where Y₁ is O, a lactate ester is formed, and where Y₁ is N(R₂), N(OR₂) or N(N(R₂)₂), a phosphoramidate ester results.

In its ester-forming role, a protecting group typically is bound to any acidic group such as, by way of example and not limitation, a -CO₂H or -C(S)OH group, thereby resulting in -CO₂R^x where R^x is defined herein. Also, R^x for example includes the enumerated ester groups of WO 95/07920.

Examples of protecting groups include:

C₃-C₁₂ heterocycle (described above) or aryl. These aromatic groups optionally are polycyclic or monocyclic. Examples include phenyl, spiryl, 2- and 3-pyrrolyl, 2- and 3-thienyl, 2- and 4-imidazolyl, 2-, 4- and 5-oxazolyl, 3- and 4-isoxazolyl, 2-, 4- and 5-thiazolyl, 3-, 4- and 5-isothiazolyl, 3- and 4-

pyrazolyl, 1-, 2-, 3- and 4-pyridinyl, and 1-, 2-, 4- and 5-pyrimidinyl,

C₃-C₁₂ heterocycle or aryl substituted with halo, R¹, R¹-O-C₁-C₁₂

alkylene, C₁-C₁₂ alkoxy, CN, NO₂, OH, carboxy, carboxyester, thiol, thioester,

C₁-C₁₂ haloalkyl (1-6 halogen atoms), C₂-C₁₂ alkenyl or C₂-C₁₂ alkynyl. Such

5 groups include 2-, 3- and 4-alkoxyphenyl (C₁-C₁₂ alkyl), 2-, 3- and 4-

methoxyphenyl, 2-, 3- and 4-ethoxyphenyl, 2,3-, 2,4-, 2,5-, 2,6-, 3,4- and 3,5-

diethoxyphenyl, 2- and 3-carboethoxy-4-hydroxyphenyl, 2- and 3-ethoxy-4-

hydroxyphenyl, 2- and 3-ethoxy-5-hydroxyphenyl, 2- and 3-ethoxy-6-

hydroxyphenyl, 2-, 3- and 4-O-acetylphenyl, 2-, 3- and 4-dimethylaminophenyl,

10 2-, 3- and 4-methylmercaptophenyl, 2-, 3- and 4-halophenyl (including 2-, 3-

and 4-fluorophenyl and 2-, 3- and 4-chlorophenyl), 2,3-, 2,4-, 2,5-, 2,6-, 3,4- and

3,5-dimethylphenyl, 2,3-, 2,4-, 2,5-, 2,6-, 3,4- and 3,5-biscarboxyethylphenyl,

2,3-, 2,4-, 2,5-, 2,6-, 3,4- and 3,5-dimethoxyphenyl, 2,3-, 2,4-, 2,5-, 2,6-, 3,4-

and 3,5-dihalophenyl (including 2,4-difluorophenyl and 3,5-difluorophenyl), 2-,

15 3- and 4-haloalkylphenyl (1 to 5 halogen atoms, C₁-C₁₂ alkyl including 4-

trifluoromethylphenyl), 2-, 3- and 4-cyanophenyl, 2-, 3- and 4-nitrophenyl, 2-,

3- and 4-haloalkylbenzyl (1 to 5 halogen atoms, C₁-C₁₂ alkyl including 4-

trifluoromethylbenzyl and 2-, 3- and 4-trichloromethylphenyl and 2-, 3- and 4-

trichloromethylphenyl), 4-N-methylpiperidinyl, 3-N-methylpiperidinyl, 1-

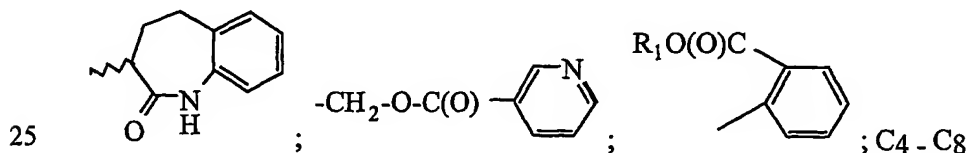
20 ethylpiperazinyl, benzyl, alkylsalicylphenyl (C₁-C₄ alkyl, including 2-, 3- and 4-

ethylsalicylphenyl), 2-, 3- and 4-acetylphenyl, 1,8-dihydroxynaphthyl (-C₁₀H₆-

OH) and aryloxy ethyl [C₆-C₉ aryl (including phenoxy ethyl)], 2,2'-

dihydroxybiphenyl, 2-, 3- and 4-N,N-dialkylaminophenol, -C₆H₄CH₂-N(CH₃)₂,

trimethoxybenzyl, triethoxybenzyl, 2-alkyl pyridinyl (C₁-4 alkyl);



esters of 2-carboxyphenyl; and C₁-C₄ alkylene-C₃-C₆ aryl (including benzyl, -

-CH₂-pyrrolyl, -CH₂-thienyl, -CH₂-imidazolyl, -CH₂-oxazolyl, -CH₂-isoxazolyl,

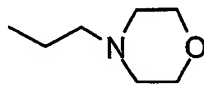
-CH₂-thiazolyl, -CH₂-isothiazolyl, -CH₂-pyrazolyl, -CH₂-pyridinyl and -CH₂-

pyrimidinyl) substituted in the aryl moiety by 3 to 5 halogen atoms or 1 to 2

30 atoms or groups selected from halogen, C₁-C₁₂ alkoxy (including methoxy and

ethoxy), cyano, nitro, OH, C₁-C₁₂ haloalkyl (1 to 6 halogen atoms; including -CH₂CCl₃), C₁-C₁₂ alkyl (including methyl and ethyl), C₂-C₁₂ alkenyl or C₂-C₁₂ alkynyl; alkoxy ethyl [C₁-C₆ alkyl including -CH₂-CH₂-O-CH₃ (methoxy ethyl)]; alkyl substituted by any of the groups set forth above for aryl, in
 5 particular OH or by 1 to 3 halo atoms (including -CH₃, -CH(CH₃)₂, -C(CH₃)₃, -CH₂CH₃, -(CH₂)₂CH₃, -(CH₂)₃CH₃, -(CH₂)₄CH₃, -(CH₂)₅CH₃, -CH₂CH₂F, -

CH₂CH₂Cl, -CH₂CF₃, and -CH₂CCl₃);



; -N-2-

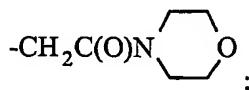
propylmorpholino, 2,3-dihydro-6-hydroxyindene, sesamol, catechol monoester, -CH₂-C(O)-N(R¹)₂, -CH₂-S(O)(R¹), -CH₂-S(O)₂(R¹), -CH₂-CH(OC(O)CH₂R¹)-
 10 CH₂(OC(O)CH₂R¹),olesteryl, enolpyruvate (HOOC-C(=CH₂)-), glycerol;
 a 5 or 6 carbon monosaccharide, disaccharide or oligosaccharide (3 to 9 monosaccharide residues);

triglycerides such as α-D-β-diglycerides (wherein the fatty acids composing glyceride lipids generally are naturally occurring saturated or
 15 unsaturated C₆₋₂₆, C₆₋₁₈ or C₆₋₁₀ fatty acids such as linoleic, lauric, myristic, palmitic, stearic, oleic, palmitoleic, linolenic and the like fatty acids) linked to acyl of the parental compounds herein through a glyceryl oxygen of the triglyceride;

phospholipids linked to the carboxyl group through the phosphate of the
 20 phospholipid;

phthalidyl (shown in Fig. 1 of Clayton et al., *Antimicrob. Agents Chemo.* (1974) 5(6):670-671);

cyclic carbonates such as (5-R_d-2-oxo-1,3-dioxolen-4-yl) methyl esters (Sakamoto et al., *Chem. Pharm. Bull.* (1984) 32(6)2241-2248) where R_d is R₁,
 25 R₄ or aryl; and



The hydroxyl groups of the compounds of this invention optionally are substituted with one of groups III, IV or V disclosed in WO 94/21604, or with isopropyl.

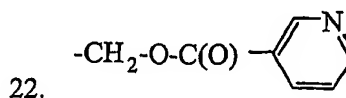
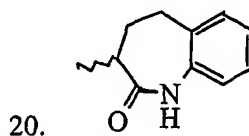
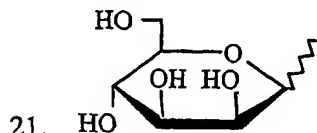
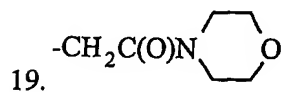
30 Table A lists examples of protecting group ester moieties that for example

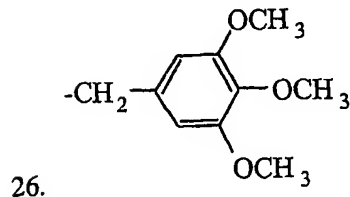
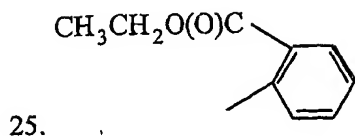
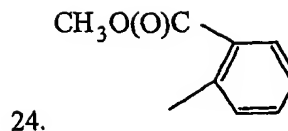
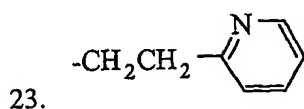
can be bonded via oxygen to $-C(O)O-$ and $-P(O)(O-)_2$ groups. Several amidates also are shown, which are bound directly to $-C(O)-$ or $-P(O)_2$. Esters of structures 1-5, 8-10 and 16, 17, 19-22 are synthesized by reacting the compound herein having a free hydroxyl with the corresponding halide (chloride or acyl chloride and the like) and N,N-dicyclohexyl-N-morpholine carboxamidate (or another base such as DBU, triethylamine, $CsCO_3$, N,N-dimethylaniline and the like) in DMF (or other solvent such as acetonitrile or N-methylpyrrolidone). When the compound to be protected is a phosphonate, the esters of structures 5-7, 11, 12, 21, and 23-26 are synthesized by reaction of the alcohol or alkoxide salt (or the corresponding amines in the case of compounds such as 13, 14 and 15) with the monochlorophosphonate or dichlorophosphonate (or another activated phosphonate).

TABLE A

15	1. $-CH_2-C(O)-N(R_1)_2^*$	10. $-CH_2-O-C(O)-C(CH_3)_3$
	2. $-CH_2-S(O)(R_1)$	11. $-CH_2-CCl_3$
	3. $-CH_2-S(O)_2(R_1)$	12. $-C_6H_5$
	4. $-CH_2-O-C(O)-CH_2-C_6H_5$	13. $-NH-CH_2-C(O)O-CH_2CH_3$
20	5. 3-cholesteryl	14. $-N(CH_3)-CH_2-C(O)O-CH_2CH_3$
	6. 3-pyridyl	15. $-NHR_1$
	7. N-ethylmorpholino	16. $-CH_2-O-C(O)-C_{10}H_{15}$
	8. $-CH_2-O-C(O)-C_6H_5$	17. $-CH_2-O-C(O)-CH(CH_3)_2$
	9. $-CH_2-O-C(O)-CH_2CH_3$	18. $-CH_2-C^*H(OC(O)CH_2R_1)-CH_2-(OC(O)CH_2R_1)^*$

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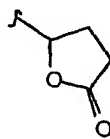




5 # - chiral center is (R), (S) or racemate.

Other esters that are suitable for use herein are described in EP 632048.

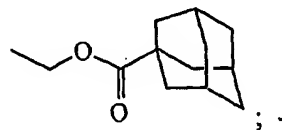
Protecting groups also includes "double ester" forming profunctionalities



such as $-\text{CH}_2\text{OC}(\text{O})\text{OCH}_3$, $-\text{CH}_2\text{SCOCCH}_3$, $-\text{CH}_2\text{OCON}(\text{CH}_3)_2$, or

10 alkyl- or aryl-acyloxyalkyl groups of the structure $-\text{CH}(\text{R}^1 \text{ or } \text{W}^5)\text{O}((\text{CO})\text{R}^{37})$ or $-\text{CH}(\text{R}^1 \text{ or } \text{W}^5)((\text{CO})\text{OR}^{38})$ (linked to oxygen of the acidic group) wherein R^{37} and R^{38} are alkyl, aryl, or alkylaryl groups (see U.S. Patent No. 4968788).

Frequently R^{37} and R^{38} are bulky groups such as branched alkyl, ortho-substituted aryl, meta-substituted aryl, or combinations thereof, including
 15 normal, secondary, iso- and tertiary alkyls of 1-6 carbon atoms. An example is the pivaloyloxymethyl group. These are of particular use with prodrugs for oral administration. Examples of such useful protecting groups are alkylacyloxymethyl esters and their derivatives, including -



$\text{CH}(\text{CH}_2\text{CH}_2\text{OCH}_3)\text{OC}(\text{O})\text{C}(\text{CH}_3)_3$,

20 $\text{CH}_2\text{OC}(\text{O})\text{C}_{10}\text{H}_{15}$, $-\text{CH}_2\text{OC}(\text{O})\text{C}(\text{CH}_3)_3$, $-\text{CH}(\text{CH}_2\text{OCH}_3)\text{OC}(\text{O})\text{C}(\text{CH}_3)_3$, -
 $\text{CH}(\text{CH}(\text{CH}_3)_2)\text{OC}(\text{O})\text{C}(\text{CH}_3)_3$, $-\text{CH}_2\text{OC}(\text{O})\text{CH}_2\text{CH}(\text{CH}_3)_2$, -
 $\text{CH}_2\text{OC}(\text{O})\text{C}_6\text{H}_{11}$, $-\text{CH}_2\text{OC}(\text{O})\text{C}_6\text{H}_5$, $-\text{CH}_2\text{OC}(\text{O})\text{C}_{10}\text{H}_{15}$, -
 $\text{CH}_2\text{OC}(\text{O})\text{CH}_2\text{CH}_3$, $-\text{CH}_2\text{OC}(\text{O})\text{CH}(\text{CH}_3)_2$, $-\text{CH}_2\text{OC}(\text{O})\text{C}(\text{CH}_3)_3$ and -
 $\text{CH}_2\text{OC}(\text{O})\text{CH}_2\text{C}_6\text{H}_5$.

In some claims the protected acidic group is an ester of the acidic group and is the residue of a hydroxyl-containing functionality. In other claims, an amino compound is used to protect the acid functionality. The residues of suitable hydroxyl or amino-containing functionalities are set forth above or are
5 found in WO 95/07920. Of particular interest are the residues of amino acids, amino acid esters, polypeptides, or aryl alcohols. Typical amino acid, polypeptide and carboxyl-esterified amino acid residues are described on pages 11-18 and related text of WO 95/07920 as groups L1 or L2. WO 95/07920 expressly teaches the amidates of phosphonic acids, but it will be understood that
10 such amidates are formed with any of the acid groups set forth herein and the amino acid residues set forth in WO 95/07920.

Typical esters for protecting acidic functionalities are also described in WO 95/07920, again understanding that the same esters can be formed with the acidic groups herein as with the phosphonate of the '920 publication. Typical
15 ester groups are defined at least on WO 95/07920 pages 89-93 (under R³¹ or R³⁵), the table on page 105, and pages 21-23 (as R). Of particular interest are esters of unsubstituted aryl such as phenyl or arylalkyl such benzyl, or hydroxy-, halo-, alkoxy-, carboxy- and/or alkylestercarboxy-substituted aryl or alkylaryl, especially phenyl, ortho-ethoxyphenyl, or C₁-C₄ alkylestercarboxyphenyl
20 (salicylate C₁-C₁₂ alkylesters).

The protected acidic groups, particularly when using the esters or amides of WO 95/07920, are useful as prodrugs for oral administration. However, it is not essential that the acidic group be protected in order for the compounds of this invention to be effectively administered by the oral route. When the compounds
25 of the invention having protected groups, in particular amino acid amidates or substituted and unsubstituted aryl esters are administered systemically or orally they are capable of hydrolytic cleavage *in vivo* to yield the free acid.

One or more of the acidic hydroxyls are protected. If more than one acidic hydroxyl is protected then the same or a different protecting group is
30 employed, e.g., the esters may be different or the same, or a mixed amidate and ester may be used.

Typical hydroxy protecting groups described in Greene (pages 14-118) include substituted methyl and alkyl ethers, substituted benzyl ethers, silyl

ethers, esters including sulfonic acid esters, and carbonates. For example:

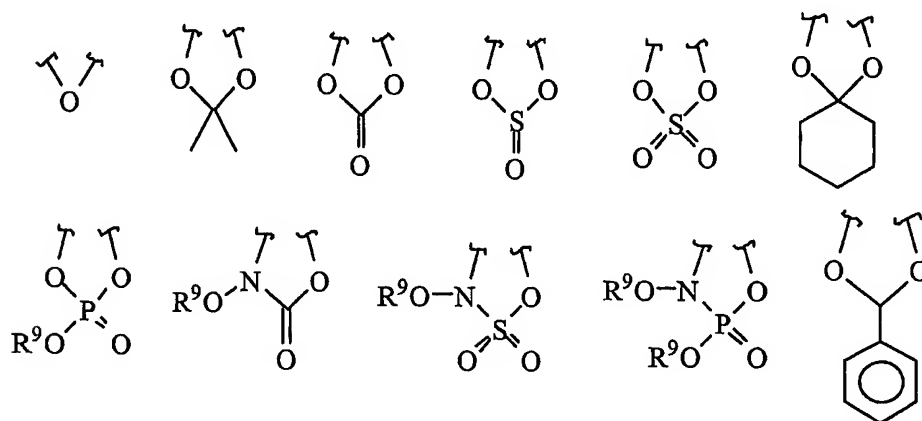
- Ethers (methyl, *t*-butyl, allyl);
- Substituted Methyl Ethers (Methoxymethyl, Methylthiomethyl, *t*-Butylthiomethyl, (Phenyldimethylsilyl)methoxymethyl, Benzyloxymethyl, *p*-Methoxybenzyloxymethyl, (4-Methoxyphenoxy)methyl, Guaiacolmethyl, *t*-Butoxymethyl, 4-Pentenylloxymethyl, Siloxymethyl, 2-Methoxyethoxymethyl, 2,2,2-Trichloroethoxymethyl, Bis(2-chloroethoxy)methyl, 2-(Trimethylsilyl)ethoxymethyl, Tetrahydropyranyl, 3-Bromotetrahydropyranyl, Tetrahydrothiopyranyl, 1-Methoxycyclohexyl, 4-Methoxytetrahydropyranyl, 4-Methoxytetrahydrothiopyranyl, 4-Methoxytetrahydrothiopyranyl *S,S*-Dioxido, 1-[(2-Chloro-4-methyl)phenyl]-4-methoxypiperidin-4-yl, 1,4-Dioxan-2-yl, Tetrahydrofuran, Tetrahydrothiofuran, 2,3,3a,4,5,6,7,7a-Octahydro-7,8,8-trimethyl-4,7-methanobenzofuran-2-yl));
- Substituted Ethyl Ethers (1-Ethoxyethyl, 1-(2-Chloroethoxy)ethyl, 1-Methyl-1-methoxyethyl, 1-Methyl-1-benzyloxyethyl, 1-Methyl-1-benzyloxy-2-fluoroethyl, 2,2,2-Trichloroethyl, 2-Trimethylsilylethyl, 2-(Phenylselenyl)ethyl, *p*-Chlorophenyl, *p*-Methoxyphenyl, 2,4-Dinitrophenyl, Benzyl);
- Substituted Benzyl Ethers (*p*-Methoxybenzyl, 3,4-Dimethoxybenzyl, *o*-Nitrobenzyl, *p*-Nitrobenzyl, *p*-Halobenzyl, 2,6-Dichlorobenzyl, *p*-Cyanobenzyl, *p*-Phenylbenzyl, 2- and 4-Picolyl, 3-Methyl-2-picolyl *N*-Oxido, Diphenylmethyl, *p,p'*-Dinitrobenzhydryl, 5-Dibenzosuberyl, Triphenylmethyl, α -Naphthylidiphenylmethyl, *p*-methoxyphenyldiphenylmethyl, Di(*p*-methoxyphenyl)phenylmethyl, Tri(*p*-methoxyphenyl)methyl, 4-(4'-Bromophenacyloxy)phenyldiphenylmethyl, 4,4',4''-Tris(4,5-dichlorophthalimidophenyl)methyl, 4,4',4''-Tris(levulinoyloxyphenyl)methyl, 4,4',4''-Tris(benzoyloxyphenyl)methyl, 3-(Imidazol-1-ylmethyl)bis(4',4''-dimethoxyphenyl)methyl, 1,1-Bis(4-methoxyphenyl)-1'-pyrenylmethyl, 9-Anthryl, 9-(9-Phenyl)xanthenyl, 9-(9-Phenyl-10-oxo)anthryl, 1,3-Benzodithiolan-2-yl, Benzisothiazolyl *S,S*-Dioxido);
- Silyl Ethers (Trimethylsilyl, Triethylsilyl, Triisopropylsilyl,

- Dimethylisopropylsilyl, Diethylisopropylsilyl, Dimethylthexylsilyl, *t*-Butyldimethylsilyl, *t*-Butyldiphenylsilyl, Tribenzylsilyl, Tri-*p*-xylylsilyl, Triphenylsilyl, Diphenylmethylsilyl, *t*-Butylmethoxyphenylsilyl);
- Esters (Formate, Benzoylformate, Acetate, Chloroacetate, Dichloroacetate, Trichloroacetate, Trifluoroacetate, Methoxyacetate, Triphenylmethoxyacetate, Phenoxyacetate, *p*-Chlorophenoxyacetate, *p*-poly-Phenylacetate, 3-Phenylpropionate, 4-Oxopentanoate (Levulinate), 4,4-(Ethylenedithio)pentanoate, Pivaloate, Adamantoate, Crotonate, 4-Methoxycrotonate, Benzoate, *p*-Phenylbenzoate, 2,4,6-Trimethylbenzoate (Mesitoate));
 - Carbonates (Methyl, 9-Fluorenylmethyl, Ethyl, 2,2,2-Trichloroethyl, 2-(Trimethylsilyl)ethyl, 2-(Phenylsulfonyl)ethyl, 2-(Triphenylphosphonio)ethyl, Isobutyl, Vinyl, Allyl, *p*-Nitrophenyl, Benzyl, *p*-Methoxybenzyl, 3,4-Dimethoxybenzyl, *o*-Nitrobenzyl, *p*-Nitrobenzyl, *S*-Benzyl Thiocarbonate, 4-Ethoxy-1-naphthyl, Methyl Dithiocarbonate);
 - Groups With Assisted Cleavage (2-Iodobenzoate, 4-Azidobutyrate, 4-Nitro-4-methylpentanoate, *o*-(Dibromomethyl)benzoate, 2-Formylbenzenesulfonate, 2-(Methylthiomethoxy)ethyl Carbonate, 4-(Methylthiomethoxy)butyrate, 2-(Methylthiomethoxymethyl)benzoate);
 - Miscellaneous Esters (2,6-Dichloro-4-methylphenoxyacetate, 2,6-Dichloro-4-(1,1,3,3 tetramethylbutyl)phenoxyacetate, 2,4-Bis(1,1-dimethylpropyl)phenoxyacetate, Chlorodiphenylacetate, Isobutyrate, Monosuccinate, (*E*)-2-Methyl-2-butenate (Tigloate), *o*-(Methoxycarbonyl)benzoate, *p*-poly-Benzoate, α -Naphthoate, Nitrate, Alkyl *N,N,N',N'*-Tetramethylphosphorodiamidate, *N*-Phenylcarbamate, Borate, Dimethylphosphinothioyl, 2,4-Dinitrophenylsulfenate); and
 - Sulfonates (Sulfate, Methanesulfonate (Mesylate), Benzylsulfonate, Tosylate).

Typical 1,2-diol protecting groups (thus, generally where two OH groups are taken together with the protecting functionality) are described in Greene at pages 118-142 and include Cyclic Acetals and Ketals (Methylene, Ethylidene, 1-*t*-Butylethylidene, 1-Phenylethylidene, (4-Methoxyphenyl)ethylidene, 2,2,2-Trichloroethylidene, Acetonide (Isopropylidene), Cyclopentylidene,

- Cyclohexylidene, Cycloheptylidene, Benzyldiene, *p*-Methoxybenzylidene, 2,4-Dimethoxybenzylidene, 3,4-Dimethoxybenzylidene, 2-Nitrobenzylidene); Cyclic Ortho Esters (Methoxymethylene, Ethoxymethylene, Dimethoxymethylene, 1-Methoxyethylidene, 1-Ethoxyethylidene, 1,2-Dimethoxyethylidene, α -Methoxybenzylidene, 1-(*N,N*-Dimethylamino)ethylidene Derivative, α -(*N,N*-Dimethylamino)benzylidene Derivative, 2-Oxacyclopentylidene); Silyl Derivatives (Di-*t*-butylsilylene Group, 1,3-(1,1,3,3-Tetraisopropylidisiloxanylidene), and Tetra-*t*-butoxydisiloxane-1,3-diylidene), Cyclic Carbonates, Cyclic Boronates, Ethyl Boronate and Phenyl Boronate.
- More typically, 1,2-diol protecting groups include those shown in Table B, still more typically, epoxides, acetonides, cyclic ketals and aryl acetals.

Table B



wherein R^9 is C_1 - C_6 alkyl.

15

Amino protecting groups

Another set of protecting groups include any of the typical amino protecting groups described by Greene at pages 315-385. They include:

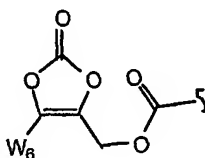
- Carbamates: (methyl and ethyl, 9-fluorenylmethyl, 9(2-sulfo)fluorenylmethyl, 9-(2,7-dibromo)fluorenylmethyl, 2,7-di-*t*-butyl-[9-(10,10-dioxo-10,10,10,10-tetrahydrothioxanthyl)]methyl, 4-methoxyphenacyl);
- Substituted Ethyl: (2,2,2-trichloroethyl, 2-trimethylsilylethyl, 2-phenylethyl, 1-(1-adamantyl)-1-methylethyl, 1,1-dimethyl-2-haloethyl, 1,1-dimethyl-2,2-

- dibromoethyl, 1,1-dimethyl-2,2,2-trichloroethyl, 1-methyl-1-(4-biphenyl)ethyl, 1-(3,5-di-*t*-butylphenyl)-1-methylethyl, 2-(2'- and 4'-pyridyl)ethyl, 2-(*N,N*-dicyclohexylcarboxamido)ethyl, *t*-butyl, 1-adamantyl, vinyl, allyl, 1-isopropylallyl, cinnamyl, 4-nitrocinnamyl, 8-quinolyl, *N*-hydroxypiperidinyl, alkylidithio, benzyl, *p*-methoxybenzyl, *p*-nitrobenzyl, *p*-bromobenzyl, *p*-chlorobenzyl, 2,4-dichlorobenzyl, 4-methylsulfinylbenzyl, 9-anthrylmethyl, diphenylmethyl);
- Groups With Assisted Cleavage: (2-methylthioethyl, 2-methylsulfonylethyl, 2-(*p*-toluenesulfonyl)ethyl, [2-(1,3-dithianyl)]methyl, 4-methylthiophenyl, 2,4-dimethylthiophenyl, 2-phosphonioethyl, 2-triphenylphosphonioisopropyl, 1,1-dimethyl-2-cyanoethyl, *m*-chloro-*p*-acyloxybenzyl, *p*-(dihydroxyboryl)benzyl, 5-benzisoxazolylmethyl, 2-(trifluoromethyl)-6-chromonylmethyl);
 - Groups Capable of Photolytic Cleavage: (*m*-nitrophenyl, 3,5-dimethoxybenzyl, *o*-nitrobenzyl, 3,4-dimethoxy-6-nitrobenzyl, phenyl(*o*-nitrophenyl)methyl); Urea-Type Derivatives (phenothiazinyl-(10)-carbonyl, *N'*-*p*-toluenesulfonylaminocarbonyl, *N'*-phenylaminothiocarbonyl);
 - Miscellaneous Carbamates: (*t*-amyl, *S*-benzyl thiocarbamate, *p*-cyanobenzyl, cyclobutyl, cyclohexyl, cyclopentyl, cyclopropylmethyl, *p*-decyloxybenzyl, diisopropylmethyl, 2,2-dimethoxycarbonylvinyl, *o*-(*N,N*-dimethylcarboxamido)benzyl, 1,1-dimethyl-3-(*N,N*-dimethylcarboxamido)propyl, 1,1-dimethylpropynyl, di(2-pyridyl)methyl, 2-furanylmethyl, 2-Iodoethyl, Isobornyl, Isobutyl, Isonicotinyl, *p*-(*p'*-Methoxyphenylazo)benzyl, 1-methylcyclobutyl, 1-methylcyclohexyl, 1-methyl-1-cyclopropylmethyl, 1-methyl-1-(3,5-dimethoxyphenyl)ethyl, 1-methyl-1-(*p*-phenylazophenyl)ethyl, 1-methyl-1-phenylethyl, 1-methyl-1-(4-pyridyl)ethyl, phenyl, *p*-(phenylazo)benzyl, 2,4,6-tri-*t*-butylphenyl, 4-(trimethylammonium)benzyl, 2,4,6-trimethylbenzyl);
 - Amides: (*N*-formyl, *N*-acetyl, *N*-chloroacetyl, *N*-trichloroacetyl, *N*-trifluoroacetyl, *N*-phenylacetyl, *N*-3-phenylpropionyl, *N*-picolinoyl, *N*-3-pyridylcarboxamide, *N*-benzoylphenylalanyl, *N*-benzoyl, *N*-*p*-phenylbenzoyl);
 - Amides With Assisted Cleavage: (*N*-*o*-nitrophenylacetyl, *N*-*o*-

- nitrophenoxyacetyl, *N*-acetoacetyl, (*N*'-dithiobenzylloxycarbonylamino)acetyl, *N*-3-(*p*-hydroxyphenyl)propionyl, *N*-3-(*o*-nitrophenyl)propionyl, *N*-2-methyl-2-(*o*-nitrophenoxy)propionyl, *N*-2-methyl-2-(*o*-phenylazophenoxy)propionyl, *N*-4-chlorobutyryl, *N*-3-methyl-3-nitrobutyryl, *N*-*o*-nitrocinnamoyl, *N*-acetylmethionine, *N*-*o*-nitrobenzoyl, *N*-*o*-(benzoyloxymethyl)benzoyl, 4,5-diphenyl-3-oxazolin-2-one);
- 5
- Cyclic Imide Derivatives: (*N*-phthalimide, *N*-dithiasuccinoyl, *N*-2,3-diphenylmaleoyl, *N*-2,5-dimethylpyrrolyl, *N*-1,1,4,4-tetramethyldisilylazacyclopentane adduct, 5-substituted 1,3-dimethyl-1,3,5-triazacyclohexan-2-one, 5-substituted 1,3-dibenzyl-1,3,5-triazacyclohexan-2-one, 1-substituted 3,5-dinitro-4-pyridonyl);
 - *N*-Alkyl and *N*-Aryl Amines: (*N*-methyl, *N*-allyl, *N*-[2-(trimethylsilyl)ethoxy]methyl, *N*-3-acetoxypropyl, *N*-(1-isopropyl-4-nitro-2-oxo-3-pyrrolin-3-yl), Quaternary Ammonium Salts, *N*-benzyl, *N*-di(4-methoxyphenyl)methyl, *N*-5-dibenzosuberyl, *N*-triphenylmethyl, *N*-(4-methoxyphenyl)diphenylmethyl, *N*-9-phenylfluorenyl, *N*-2,7-dichloro-9-fluorenylmethylene, *N*-ferrocenylmethyl, *N*-2-picolylamine *N*'-oxide);
 - Imine Derivatives: (*N*-1,1-dimethylthiomethylene, *N*-benzylidene, *N*-*p*-methoxybenzylidene, *N*-diphenylmethylene, *N*-[(2-pyridyl)mesityl]methylene, *N*-(*N*',*N*'-dimethylaminomethylene, *N*-(*N*'-isopropylidene, *N*-*p*-nitrobenzylidene, *N*-salicylidene, *N*-5-chlorosalicylidene, *N*-(5-chloro-2-hydroxyphenyl)phenylmethylene, *N*-cyclohexylidene);
 - Enamine Derivatives: (*N*-(5,5-dimethyl-3-oxo-1-cyclohexenyl));
 - *N*-Metal Derivatives (*N*-borane derivatives, *N*-diphenylborinic acid derivatives, *N*-[phenyl(pentacarbonylchromium- or -tungsten)]carbenyl, *N*-copper or *N*-zinc chelate);
 - *N*-N Derivatives: (*N*-nitro, *N*-nitroso, *N*-oxide);
 - *N*-P Derivatives: (*N*-diphenylphosphinyl, *N*-dimethylthiophosphinyl, *N*-diphenylthiophosphinyl, *N*-dialkyl phosphoryl, *N*-dibenzyl phosphoryl, *N*-diphenyl phosphoryl);
 - *N*-Si Derivatives, *N*-S Derivatives, and *N*-Sulfenyl Derivatives: (*N*-benzenesulfenyl, *N*-*o*-nitrobenzenesulfenyl, *N*-2,4-dinitrobenzenesulfenyl, *N*-pentachlorobenzenesulfenyl, *N*-2-nitro-4-methoxybenzenesulfenyl, *N*-
- 10
- 15
- 20
- 25
- 30

triphenylmethylsulfenyl, *N*-3-nitropyridinesulfenyl); and *N*-sulfenyl
 Derivatives (*N*-*p*-toluenesulfenyl, *N*-benzenesulfenyl, *N*-2,3,6-trimethyl-4-
 methoxybenzenesulfenyl, *N*-2,4,6-trimethoxybenzenesulfenyl, *N*-2,6-
 dimethyl-4-methoxybenzenesulfenyl, *N*-pentamethylbenzenesulfenyl, *N*-
 5 2,3,5,6,-tetramethyl-4-methoxybenzenesulfenyl, *N*-4-
 methoxybenzenesulfenyl, *N*-2,4,6-trimethylbenzenesulfenyl, *N*-2,6-
 dimethoxy-4-methylbenzenesulfenyl, *N*-2,2,5,7,8-pentamethylchroman-6-
 sulfenyl, *N*-methanesulfenyl, *N*- β -trimethylsilyethanesulfenyl, *N*-9-
 anthracenesulfenyl, *N*-4-(4',8'-dimethoxynaphthylmethyl)benzenesulfenyl,
 10 *N*-benzylsulfenyl, *N*-trifluoromethylsulfenyl, *N*-phenacylsulfenyl).

More typically, protected amino groups include carbamates and amides,
 still more typically, -NHC(O)R^1 or $\text{-N=CR}^1\text{N(R}^1)_2$. Another protecting group,
 also useful as a prodrug for amino or $\text{-NH(R}^5\text{)}$, is:



15 See for example Alexander, J. et al. (1996) *J. Med. Chem.* 39:480-486.

Amino acid and polypeptide protecting group and conjugates

An amino acid or polypeptide protecting group of a compound of the
 invention has the structure $\text{R}^{15}\text{NHCH(R}^{16}\text{)C(O)-}$, where R^{15} is H, an amino acid
 or polypeptide residue, or R^5 , and R^{16} is defined below.

20 R^{16} is lower alkyl or lower alkyl ($\text{C}_1\text{-C}_6$) substituted with amino,
 carboxyl, amide, carboxyl ester, hydroxyl, $\text{C}_6\text{-C}_7$ aryl, guanidiny, imidazolyl,
 indolyl, sulfhydryl, sulfoxide, and/or alkylphosphate. R^{10} also is taken together
 with the amino acid α N to form a proline residue ($\text{R}^{10} = \text{-CH}_2\text{)}_3\text{-}$). However,
 R^{10} is generally the side group of a naturally-occurring amino acid such as H, -
 25 CH_3 , $\text{-CH(CH}_3\text{)}_2$, $\text{-CH}_2\text{-CH(CH}_3\text{)}_2$, $\text{-CHCH}_3\text{-CH}_2\text{-CH}_3$, $\text{-CH}_2\text{-C}_6\text{H}_5$, $\text{-CH}_2\text{CH}_2\text{-}$
 S-CH_3 , $\text{-CH}_2\text{OH}$, -CH(OH)-CH_3 , $\text{-CH}_2\text{-SH}$, $\text{-CH}_2\text{-C}_6\text{H}_4\text{OH}$, $\text{-CH}_2\text{-CO-NH}_2$, -
 $\text{CH}_2\text{-CH}_2\text{-CO-NH}_2$, $\text{-CH}_2\text{-COOH}$, $\text{-CH}_2\text{-CH}_2\text{-COOH}$, $\text{-(CH}_2\text{)}_4\text{-NH}_2$ and -
 $\text{(CH}_2\text{)}_3\text{-NH-C(NH}_2\text{)-NH}_2$. R_{10} also includes 1-guanidinoprop-3-yl, benzyl, 4-
 hydroxybenzyl, imidazol-4-yl, indol-3-yl, methoxyphenyl and ethoxyphenyl.

Another set of protecting groups include the residue of an amino-containing compound, in particular an amino acid, a polypeptide, a protecting group, $-\text{NH}\text{SO}_2\text{R}$, $\text{NHC}(\text{O})\text{R}$, $-\text{N}(\text{R})_2$, NH_2 or $-\text{NH}(\text{R})(\text{H})$, whereby for example a carboxylic acid is reacted, i.e. coupled, with the amine to form an amide, as in
5 $\text{C}(\text{O})\text{NR}_2$. A phosphonic acid may be reacted with the amine to form a phosphoramidate, as in $-\text{P}(\text{O})(\text{OR})(\text{NR}_2)$.

In general, amino acids have the structure $\text{R}^{17}\text{C}(\text{O})\text{CH}(\text{R}^{16})\text{NH}-$, where R^{17} is $-\text{OH}$, $-\text{OR}$, an amino acid or a polypeptide residue. Amino acids are low molecular weight compounds, on the order of less than about 1000 MW and
10 which contain at least one amino or imino group and at least one carboxyl group. Generally the amino acids will be found in nature, i.e., can be detected in biological material such as bacteria or other microbes, plants, animals or man. Suitable amino acids typically are alpha amino acids, i.e. compounds characterized by one amino or imino nitrogen atom separated from the carbon
15 atom of one carboxyl group by a single substituted or unsubstituted alpha carbon atom. Of particular interest are hydrophobic residues such as mono- or di-alkyl or aryl amino acids, cycloalkylamino acids and the like. These residues contribute to cell permeability by increasing the partition coefficient of the parental drug. Typically, the residue does not contain a sulfhydryl or guanidino
20 substituent.

Naturally-occurring amino acid residues are those residues found naturally in plants, animals or microbes, especially proteins thereof. Polypeptides most typically will be substantially composed of such naturally-occurring amino acid residues. These amino acids are glycine, alanine, valine,
25 leucine, isoleucine, serine, threonine, cysteine, methionine, glutamic acid, aspartic acid, lysine, hydroxylysine, arginine, histidine, phenylalanine, tyrosine, tryptophan, proline, asparagine, glutamine and hydroxyproline. Additionally, unnatural amino acids, for example, valanine, phenylglycine and homoarginine are also included. Commonly encountered amino acids that are not gene-
30 encoded may also be used in the present invention. All of the amino acids used in the present invention may be either the D- or L- optical isomer. In addition, other peptidomimetics are also useful in the present invention. For a general review, see Spatola, A. F., in Chemistry and Biochemistry of Amino Acids,

Peptides and Proteins, B. Weinstein, eds., Marcel Dekker, New York, p. 267 (1983).

When protecting groups are single amino acid residues or polypeptides they optionally are substituted at R^3 of substituents A^1 , A^2 or A^3 in Formula I.

- 5 These conjugates are produced by forming an amide bond between a carboxyl group of the amino acid (or C-terminal amino acid of a polypeptide for example). Similarly, conjugates are formed between R^3 (Formula I) and an amino group of an amino acid or polypeptide. Generally, only one of any site in the parental molecule is amidated with an amino acid as described herein,
- 10 although it is within the scope of this invention to introduce amino acids at more than one permitted site. Usually, a carboxyl group of R^3 is amidated with an amino acid. In general, the α -amino or α -carboxyl group of the amino acid or the terminal amino or carboxyl group of a polypeptide are bonded to the parental functionalities, i.e., carboxyl or amino groups in the amino acid side chains
- 15 generally are not used to form the amide bonds with the parental compound (although these groups may need to be protected during synthesis of the conjugates as described further below).

- With respect to the carboxyl-containing side chains of amino acids or polypeptides it will be understood that the carboxyl group optionally will be
- 20 blocked, e.g., by R^1 , esterified with R^5 or amidated. Similarly, the amino side chains R^{16} optionally will be blocked with R^1 or substituted with R^5 .

Such ester or amide bonds with side chain amino or carboxyl groups, like the esters or amides with the parental molecule, optionally are hydrolyzable *in vivo* or *in vitro* under acidic (pH <3) or basic (pH >10) conditions.

- 25 Alternatively, they are substantially stable in the gastrointestinal tract of humans but are hydrolyzed enzymatically in blood or in intracellular environments. The esters or amino acid or polypeptide amidates also are useful as intermediates for the preparation of the parental molecule containing free amino or carboxyl groups. The free acid or base of the parental compound, for example, is readily
- 30 formed from the esters or amino acid or polypeptide conjugates of this invention by conventional hydrolysis procedures.

When an amino acid residue contains one or more chiral centers, any of the D, L, meso, threo or erythro (as appropriate) racemates, scalemates or

mixtures thereof may be used. In general, if the intermediates are to be hydrolyzed non-enzymatically (as would be the case where the amides are used as chemical intermediates for the free acids or free amines), D isomers are useful. On the other hand, L isomers are more versatile since they can be susceptible to both non-enzymatic and enzymatic hydrolysis, and are more efficiently transported by amino acid or dipeptidyl transport systems in the gastrointestinal tract.

Examples of suitable amino acids whose residues are represented by R^x or R^y include the following:

10 Glycine;

Aminopolycarboxylic acids, e.g., aspartic acid, β -hydroxyaspartic acid, glutamic acid, β -hydroxyglutamic acid, β -methylaspartic acid, β -methylglutamic acid, β , β -dimethylaspartic acid, γ -hydroxyglutamic acid, β , γ -dihydroxyglutamic acid, β -phenylglutamic acid, γ -methyleneglutamic acid, 3-aminoadipic acid, 2-aminopimelic acid, 2-aminosuberic acid and 2-aminosebacic acid;

15 Amino acid amides such as glutamine and asparagine;

Polyamino- or polybasic-monocarboxylic acids such as arginine, lysine, β -aminoalanine, γ -aminobutyrate, ornithine, citrulline, homoarginine, homocitrulline, hydroxylysine, alcohohydroxylysine and diaminobutyric acid;

20 Other basic amino acid residues such as histidine;

Diaminodicarboxylic acids such as α , α' -diaminosuccinic acid, α , α' -diaminoglutaric acid, α , α' -diaminoadipic acid, α , α' -diaminopimelic acid, α , α' -diamino- β -hydroxypimelic acid, α , α' -diaminosuberic acid, α , α' -diaminoazelaic acid, and α , α' -diaminosebacic acid;

25 Imino acids such as proline, hydroxyproline, alcohohydroxyproline, γ -methylproline, pipercolic acid, 5-hydroxypipercolic acid, and azetidine-2-carboxylic acid;

A mono- or di-alkyl (typically C₁-C₈ branched or normal) amino acid such as alanine, valine, leucine, allylglycine, butyrate, norvaline, norleucine, 30 heptyline, α -methylserine, α -amino- α -methyl- γ -hydroxyvaleric acid, α -amino- α -methyl- δ -hydroxyvaleric acid, α -amino- α -methyl- ϵ -hydroxycaproic acid, isovaline, α -methylglutamic acid, α -aminoisobutyric acid, α -aminodiethylacetic acid, α -aminodiisopropylacetic acid, α -aminodi-n-propylacetic acid, α -

- aminodiisobutylic acid, α -aminodi-n-butylic acid, α -aminoethylisopropylacetic acid, α -amino-n-propylacetic acid, α -aminodiisoamyacetic acid, α -methylaspartic acid, α -methylglutamic acid, 1-aminocyclopropane-1-carboxylic acid, isoleucine, alloisoleucine, *tert*-leucine, β -methyltryptophan and α -amino- β -ethyl- β -phenylpropionic acid;
- 5 β -phenylserinyl;
- Aliphatic α -amino- β -hydroxy acids such as serine, β -hydroxyleucine, β -hydroxynorleucine, β -hydroxynorvaline, and α -amino- β -hydroxystearic acid;
- α -Amino, α -, γ -, δ - or ϵ -hydroxy acids such as homoserine, δ -
- 10 hydroxynorvaline, γ -hydroxynorvaline and ϵ -hydroxynorleucine residues;
- canavine and canaline; γ -hydroxyornithine;
- 2-hexosaminic acids such as D-glucosaminic acid or D-galactosaminic acid;
- α -Amino- β -thiols such as penicillamine, β -thiolnorvaline or β -
- 15 thiolbutyrine;
- Other sulfur containing amino acid residues including cysteine;
- homocystine, β -phenylmethionine, methionine, S-allyl-L-cysteine sulfoxide, 2-thiolhistidine, cystathionine, and thiol ethers of cysteine or homocysteine;
- Phenylalanine, tryptophan and ring-substituted α -amino acids such as the
- 20 phenyl- or cyclohexylamino acids α -aminophenylacetic acid, α -aminocyclohexylacetic acid and α -amino- β -cyclohexylpropionic acid;
- phenylalanine analogues and derivatives comprising aryl, lower alkyl, hydroxy, guanidino, oxyalkylether, nitro, sulfur or halo-substituted phenyl (e.g., tyrosine, methyltyrosine and o-chloro-, p-chloro-, 3,4-dichloro, *o*-, *m*- or *p*-methyl-, 2,4,6-
- 25 trimethyl-, 2-ethoxy-5-nitro-, 2-hydroxy-5-nitro- and p-nitro-phenylalanine);
- furyl-, thienyl-, pyridyl-, pyrimidinyl-, purinyl- or naphthyl-alanines; and
- tryptophan analogues and derivatives including kynurenine, 3-hydroxykynurenine, 2-hydroxytryptophan and 4-carboxytryptophan;
- α -Amino substituted amino acids including sarcosine (N-methylglycine),
- 30 N-benzylglycine, N-methylalanine, N-benzylalanine, N-methylphenylalanine, N-benzylphenylalanine, N-methylvaline and N-benzylvaline; and
- α -Hydroxy and substituted α -hydroxy amino acids including serine, threonine, allothreonine, phosphoserine and phosphothreonine.

Polypeptides are polymers of amino acids in which a carboxyl group of one amino acid monomer is bonded to an amino or imino group of the next amino acid monomer by an amide bond. Polypeptides include dipeptides, low molecular weight polypeptides (about 1500-5000 MW) and proteins. Proteins optionally contain 3, 5, 10, 50, 75, 100 or more residues, and suitably are substantially sequence-homologous with human, animal, plant or microbial proteins. They include enzymes (e.g., hydrogen peroxidase) as well as immunogens such as KLH, or antibodies or proteins of any type against which one wishes to raise an immune response. The nature and identity of the polypeptide may vary widely.

The polypeptide amidates are useful as immunogens in raising antibodies against either the polypeptide (if it is not immunogenic in the animal to which it is administered) or against the epitopes on the remainder of the compound of this invention.

Antibodies capable of binding to the parental non-peptidyl compound are used to separate the parental compound from mixtures, for example in diagnosis or manufacturing of the parental compound. The conjugates of parental compound and polypeptide generally are more immunogenic than the polypeptides in closely homologous animals, and therefore make the polypeptide more immunogenic for facilitating raising antibodies against it. Accordingly, the polypeptide or protein may not need to be immunogenic in an animal typically used to raise antibodies, e.g., rabbit, mouse, horse, or rat, but the final product conjugate should be immunogenic in at least one of such animals. The polypeptide optionally contains a peptidolytic enzyme cleavage site at the peptide bond between the first and second residues adjacent to the acidic heteroatom. Such cleavage sites are flanked by enzymatic recognition structures, e.g., a particular sequence of residues recognized by a peptidolytic enzyme.

Peptidolytic enzymes for cleaving the polypeptide conjugates of this invention are well known, and in particular include carboxypeptidases. Carboxypeptidases digest polypeptides by removing C-terminal residues, and are specific in many instances for particular C-terminal sequences. Such enzymes and their substrate requirements in general are well known. For example, a

dipeptide (having a given pair of residues and a free carboxyl terminus) is covalently bonded through its α -amino group to the phosphorus or carbon atoms of the compounds herein. In claims where W_1 is phosphonate it is expected that this peptide will be cleaved by the appropriate peptidolytic enzyme, leaving the
 5 carboxyl of the proximal amino acid residue to autocatalytically cleave the phosphonoamidate bond.

Suitable dipeptidyl groups (designated by their single letter code) are
 AA, AR, AN, AD, AC, AE, AQ, AG, AH, AI, AL, AK, AM, AF, AP, AS, AT,
 AW, AY, AV, RA, RR, RN, RD, RC, RE, RQ, RG, RH, RI, RL, RK, RM, RF,
 10 RP, RS, RT, RW, RY, RV, NA, NR, NN, ND, NC, NE, NQ, NG, NH, NI, NL,
 NK, NM, NF, NP, NS, NT, NW, NY, NV, DA, DR, DN, DD, DC, DE, DQ, DG,
 DH, DI, DL, DK, DM, DF, DP, DS, DT, DW, DY, DV, CA, CR, CN, CD, CC,
 CE, CQ, CG, CH, CI, CL, CK, CM, CF, CP, CS, CT, CW, CY, CV, EA, ER,
 EN, ED, EC, EE, EQ, EG, EH, EI, EL, EK, EM, EF, EP, ES, ET, EW, EY, EV,
 15 QA, QR, QN, QD, QC, QE, QQ, QG, QH, QI, QL, QK, QM, QF, QP, QS, QT,
 QW, QY, QV, GA, GR, GN, GD, GC, GE, GQ, GG, GH, GI, GL, GK, GM, GF,
 GP, GS, GT, GW, GY, GV, HA, HR, HN, HD, HC, HE, HQ, HG, HH, HI, HL,
 HK, HM, HF, HP, HS, HT, HW, HY, HV, IA, IR, IN, ID, IC, IE, IQ, IG, IH, IL,
 IL, IK, IM, IF, IP, IS, IT, IW, IY, IV, LA, LR, LN, LD, LC, LE, LQ, LG, LH,
 20 LI, LL, LK, LM, LF, LP, LS, LT, LW, LY, LV, KA, KR, KN, KD, KC, KE,
 KQ, KG, KH, KI, KL, KK, KM, KF, KP, KS, KT, KW, KY, KV, MA, MR,
 MN, MD, MC, ME, MQ, MG, MH, MI, ML, MK, MM, MF, MP, MS, MT,
 MW, MY, MV, FA, FR, FN, FD, FC, FE, FQ, FG, FH, FI, FL, FK, FM, FF, FP,
 FS, FT, FW, FY, FV, PA, PR, PN, PD, PC, PE, PQ, PG, PH, PI, PL, PK, PM,
 25 PF, PP, PS, PT, PW, PY, PV, SA, SR, SN, SD, SC, SE, SQ, SG, SH, SI, SL, SK,
 SM, SF, SP, SS, ST, SW, SY, SV, TA, TR, TN, TD, TC, TE, TQ, TG, TH, TI,
 TL, TK, TM, TF, TP, TS, TT, TW, TY, TV, WA, WR, WN, WD, WC, WE,
 WQ, WG, WH, WI, WL, WK, WM, WF, WP, WS, WT, WW, WY, WV, YA,
 YR, YN, YD, YC, YE, YQ, YG, YH, YI, YL, YK, YM, YF, YP, YS, YT, YW,
 30 YY, YV, VA, VR, VN, VD, VC, VE, VQ, VG, VH, VI, VL, VK, VM, VF, VP,
 VS, VT, VW, VY and VV.

Tripeptide residues are also useful as protecting groups. When a phosphonate is to be protected, the sequence $-X^4\text{-pro-X}^5-$ (where X^4 is any amino

acid residue and X⁵ is an amino acid residue, a carboxyl ester of proline, or hydrogen) will be cleaved by luminal carboxypeptidase to yield X⁴ with a free carboxyl, which in turn is expected to autocatalytically cleave the phosphonoamidate bond. The carboxy group of X⁵ optionally is esterified with
5 benzyl.

Dipeptide or tripeptide species can be selected on the basis of known transport properties and/or susceptibility to peptidases that can affect transport to intestinal mucosal or other cell types. Dipeptides and tripeptides lacking an α -amino group are transport substrates for the peptide transporter found in brush
10 border membrane of intestinal mucosal cells (Bai, J.P.F., (1992) *Pharm Res.* 9:969-978). Transport competent peptides can thus be used to enhance bioavailability of the amidate compounds. Di- or tripeptides having one or more amino acids in the D configuration are also compatible with peptide transport and can be utilized in the amidate compounds of this invention. Amino acids in
15 the D configuration can be used to reduce the susceptibility of a di- or tripeptide to hydrolysis by proteases common to the brush border such as aminopeptidase N. In addition, di- or tripeptides alternatively are selected on the basis of their relative resistance to hydrolysis by proteases found in the lumen of the intestine. For example, tripeptides or polypeptides lacking asp and/or glu are poor
20 substrates for aminopeptidase A, di- or tripeptides lacking amino acid residues on the N-terminal side of hydrophobic amino acids (leu, tyr, phe, val, trp) are poor substrates for endopeptidase, and peptides lacking a pro residue at the penultimate position at a free carboxyl terminus are poor substrates for carboxypeptidase P. Similar considerations can also be applied to the selection
25 of peptides that are either relatively resistant or relatively susceptible to hydrolysis by cytosolic, renal, hepatic, serum or other peptidases. Such poorly cleaved polypeptide amidates are immunogens or are useful for bonding to proteins in order to prepare immunogens.

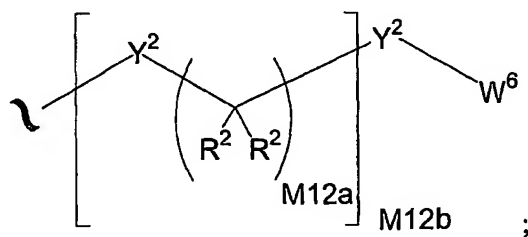
30 Specific Embodiments of the Invention

Specific values described for radicals, substituents, and ranges, as well as specific embodiments of the invention described herein, are for illustration only; they do not exclude other defined values or other values within defined ranges.

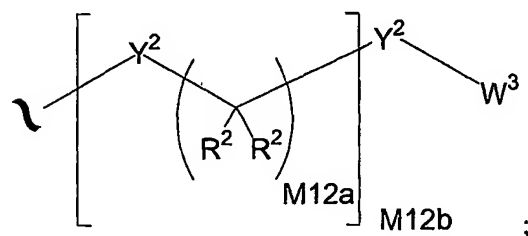
In one specific embodiment of the invention, the conjugate is a compound that is substituted with one or more phosphonate groups either directly or indirectly through a linker; and that is optionally substituted with one or more groups A^0 ; or a pharmaceutically acceptable salt thereof, wherein:

5 A^0 is A^1 , A^2 or W^3 ;

A^1 is:

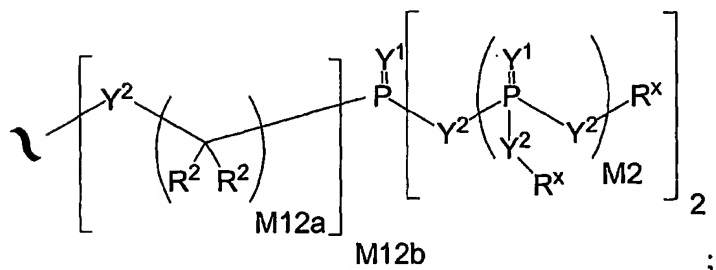


A^2 is:



10

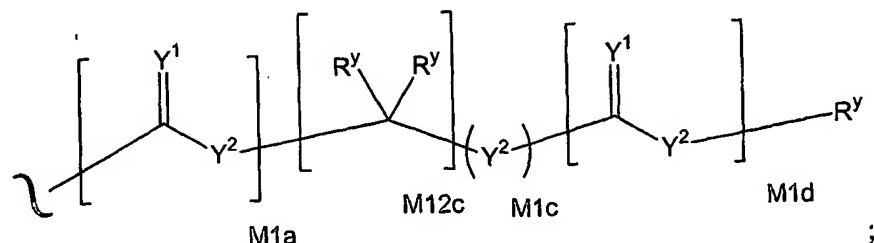
A^3 is:



Y^1 is independently O, S, $N(R^x)$, $N(O)(R^x)$, $N(OR^x)$, $N(O)(OR^x)$, or $N(N(R^x))(R^x)$;

15 Y^2 is independently a bond, O, $N(R^x)$, $N(O)(R^x)$, $N(OR^x)$, $N(O)(OR^x)$, $N(N(R^x))(R^x)$, $-S(O)_{M2-}$, or $-S(O)_{M2-}S(O)_{M2-}$;

R^x is independently H, R^1 , W^3 , a protecting group, or the formula:



wherein:

R^y is independently H, W^3 , R^2 or a protecting group;

R^1 is independently H or alkyl of 1 to 18 carbon atoms;

5 R^2 is independently H, R^1 , R^3 or R^4 wherein each R^4 is independently substituted with 0 to 3 R^3 groups or taken together at a carbon atom, two R^2 groups form a ring of 3 to 8 carbons and the ring may be substituted with 0 to 3 R^3 groups;

10 R^3 is R^{3a} , R^{3b} , R^{3c} or R^{3d} , provided that when R^3 is bound to a heteroatom, then R^3 is R^{3c} or R^{3d} ;

R^{3a} is F, Cl, Br, I, -CN, N_3 or -NO₂;

R^{3b} is Y^1 ;

15 R^{3c} is - R^x , - $N(R^x)(R^x)$, - SR^x , - $S(O)R^x$, - $S(O)_2R^x$, - $S(O)(OR^x)$, - $S(O)_2(OR^x)$, - $OC(Y^1)R^x$, - $OC(Y^1)OR^x$, - $OC(Y^1)(N(R^x)(R^x))$, - $SC(Y^1)R^x$, - $SC(Y^1)OR^x$, - $SC(Y^1)(N(R^x)(R^x))$, - $N(R^x)C(Y^1)R^x$, - $N(R^x)C(Y^1)OR^x$, or - $N(R^x)C(Y^1)(N(R^x)(R^x))$;

R^{3d} is - $C(Y^1)R^x$, - $C(Y^1)OR^x$ or - $C(Y^1)(N(R^x)(R^x))$;

R^4 is an alkyl of 1 to 18 carbon atoms, alkenyl of 2 to 18 carbon atoms, or alkynyl of 2 to 18 carbon atoms;

20 R^5 is R^4 wherein each R^4 is substituted with 0 to 3 R^3 groups;

R^{5a} is independently alkylene of 1 to 18 carbon atoms, alkenylene of 2 to 18 carbon atoms, or alkynylene of 2-18 carbon atoms any one of which alkylene, alkenylene or alkynylene is substituted with 0-3 R^3 groups;

W^3 is W^4 or W^5 ;

25 W^4 is R^5 , - $C(Y^1)R^5$, - $C(Y^1)W^5$, -SO₂ R^5 , or -SO₂ W^5 ;

W^5 is carbocycle or heterocycle wherein W^5 is independently substituted with 0 to 3 R^2 groups;

W^6 is W^3 independently substituted with 1, 2, or 3 A^3 groups;

M2 is 0, 1 or 2;

M12a is 1, 2, 3, 4, 5, 6, 7, 8, 9, 10, 11 or 12;

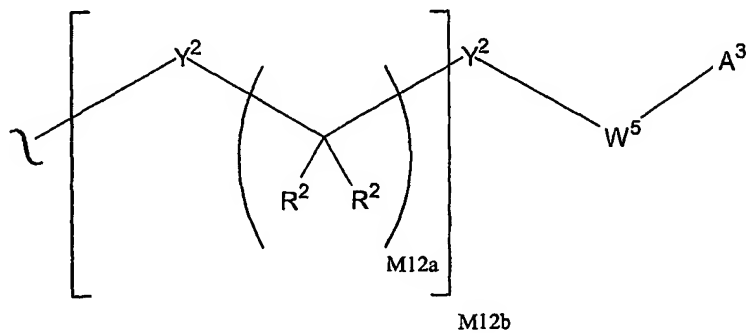
M12b is 0, 1, 2, 3, 4, 5, 6, 7, 8, 9, 10, 11 or 12;

M1a, M1c, and M1d are independently 0 or 1; and

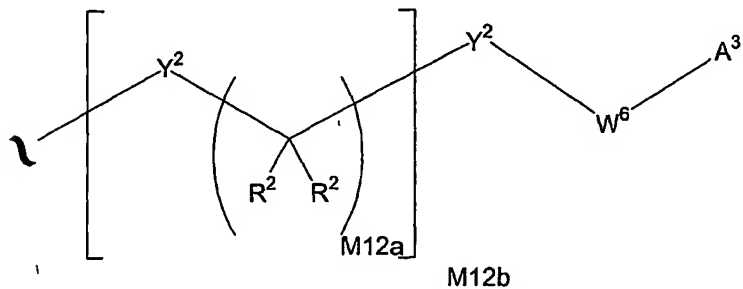
M12c is 0, 1, 2, 3, 4, 5, 6, 7, 8, 9, 10, 11 or 12.

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In another specific embodiment of the invention A¹ is of the formula:

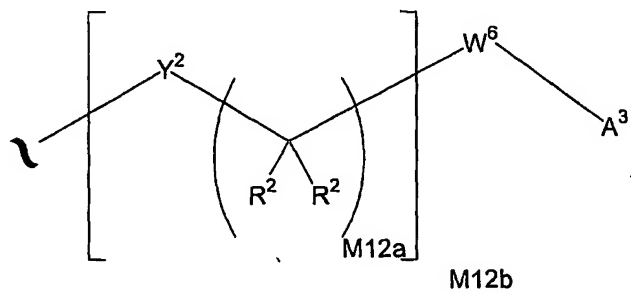


In another specific embodiment of the invention A¹ is of the formula:

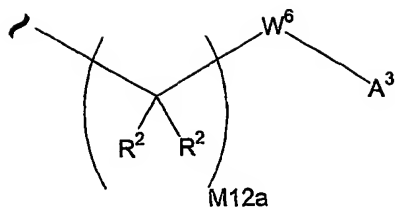


10

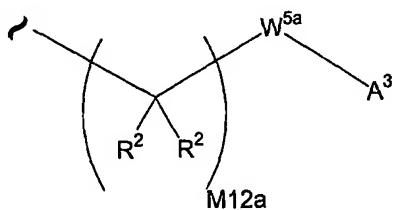
In another specific embodiment of the invention A¹ is of the formula:



In another specific embodiment of the invention A¹ is of the formula:



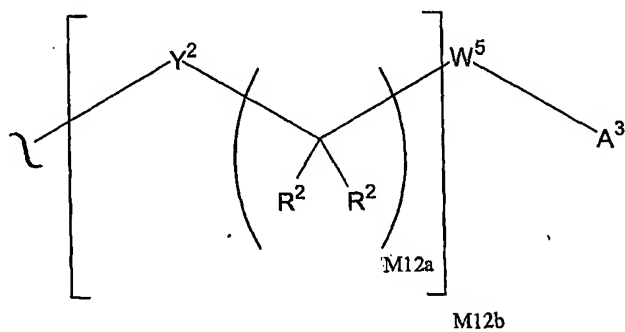
In another specific embodiment of the invention A^1 is of the formula:



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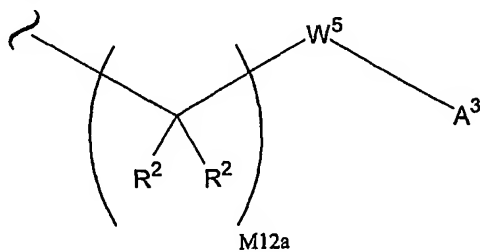
and W^{5a} is a carbocycle or a heterocycle where W^{5a} is independently substituted with 0 or 1 R^2 groups. A specific value for M12a is 1.

In another specific embodiment of the invention A^1 is of the formula:



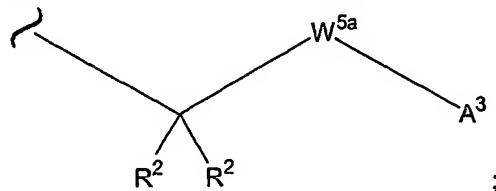
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In another specific embodiment of the invention A^1 is of the formula:



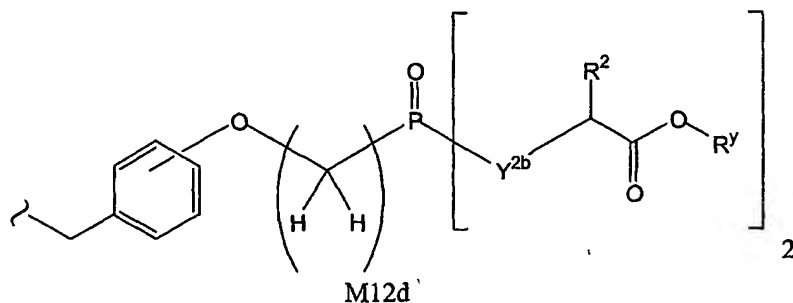
15

In another specific embodiment of the invention A^1 is of the formula:



wherein W^{5a} is a carbocycle independently substituted with 0 or 1 R^2 groups;

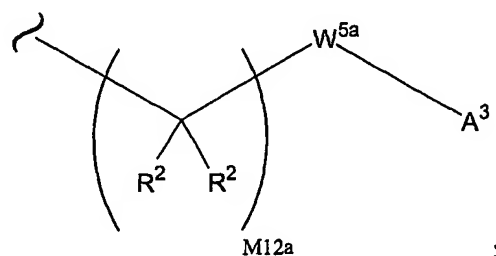
5 In another specific embodiment of the invention A^1 is of the formula:



wherein Y^{2b} is O or $N(R^2)$; and M12d is 1, 2, 3, 4, 5, 6, 7 or 8.

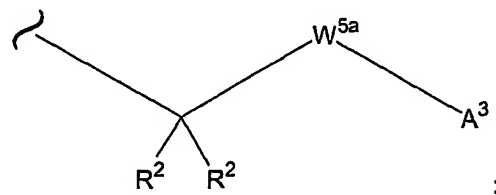
In another specific embodiment of the invention A^1 is of the formula:

10



wherein W^{5a} is a carbocycle independently substituted with 0 or 1 R^2 groups;

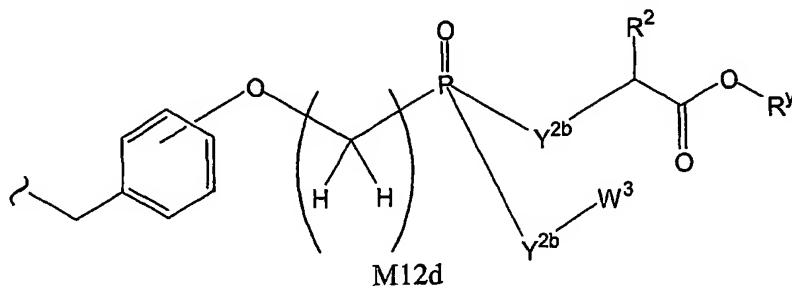
In another specific embodiment of the invention A^1 is of the formula:



15

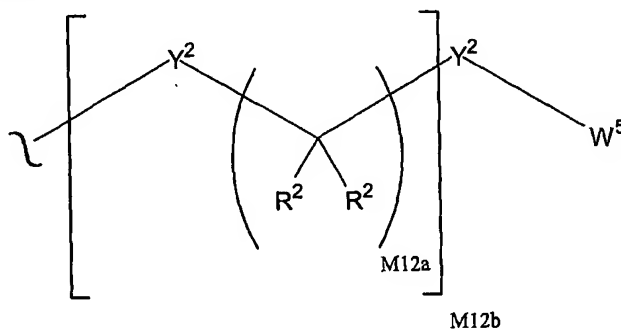
wherein W^{5a} is a carbocycle or heterocycle where W^{5a} is independently substituted with 0 or 1 R^2 groups.

In another specific embodiment of the invention A¹ is of the formula:

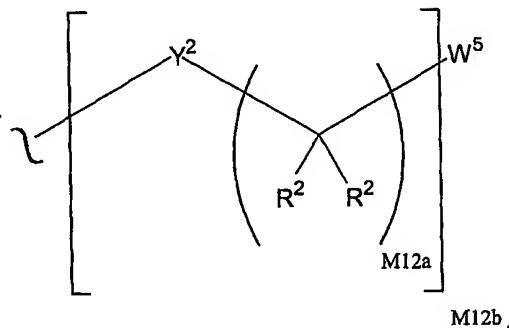


wherein Y^{2b} is O or N(R²); and M12d is 1, 2, 3, 4, 5, 6, 7 or 8.

5 In a specific embodiment of the invention A² is of the formula:



In another specific embodiment of the invention A² is of the formula:



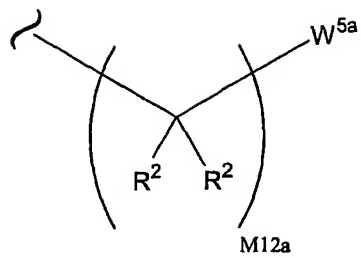
10

In another specific embodiment of the invention M12b is 1.

In another specific embodiment of the invention M12b is 0, Y² is a bond and W⁵ is a carbocycle or heterocycle where W⁵ is optionally and independently substituted with 1, 2, or 3 R² groups.

15

In another specific embodiment of the invention A² is of the formula:

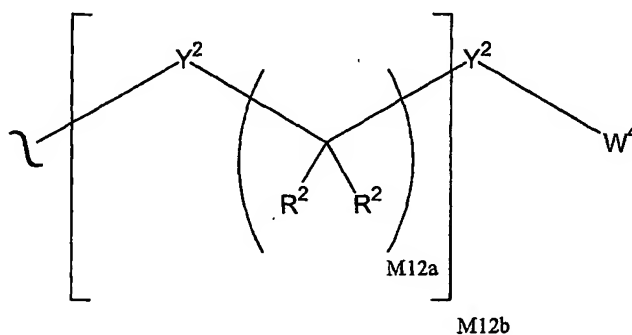


wherein W^{5a} is a carbocycle or heterocycle where W^{5a} is optionally and independently substituted with 1, 2, or 3 R^2 groups.

In another specific embodiment of the invention M12a is 1.

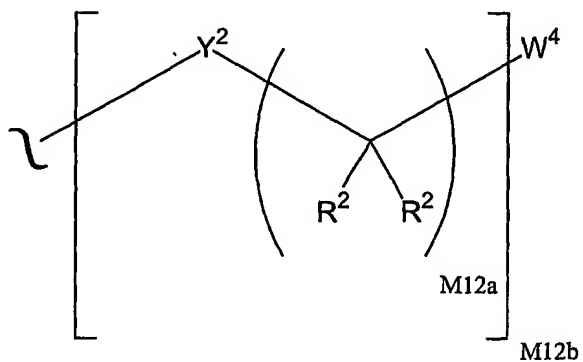
- 5 In another specific embodiment of the invention A^2 is selected from phenyl, substituted phenyl, benzyl, substituted benzyl, pyridyl and substituted pyridyl.

In another specific embodiment of the invention A^2 is of the formula:



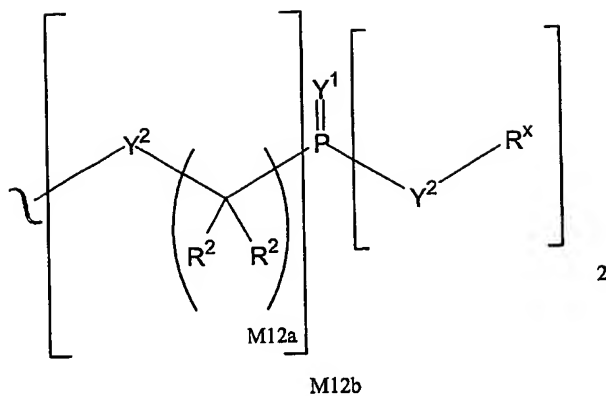
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In another specific embodiment of the invention A^2 is of the formula:

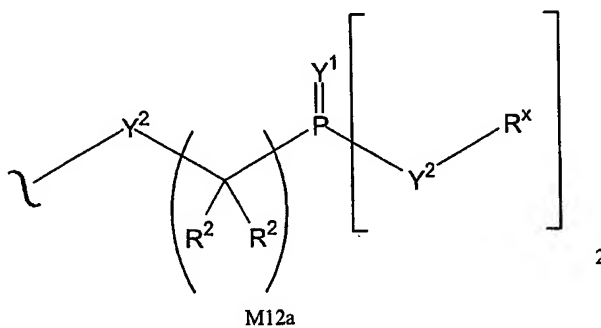


In another specific embodiment of the invention M12b is 1.

- 15 In a specific embodiment of the invention A^3 is of the formula:

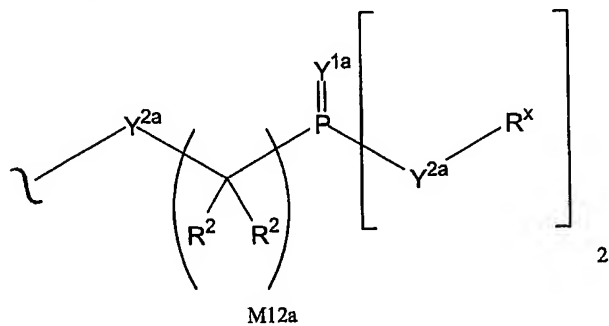


In another specific embodiment of the invention A³ is of the formula:



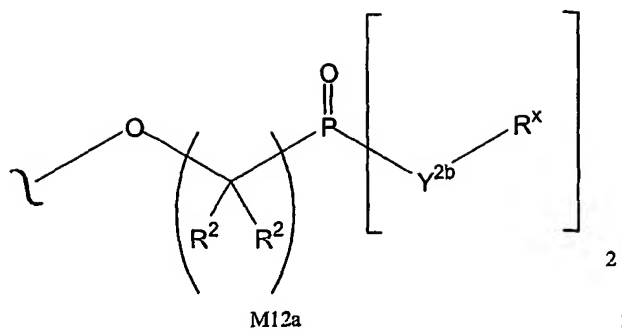
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In another specific embodiment of the invention A³ is of the formula:



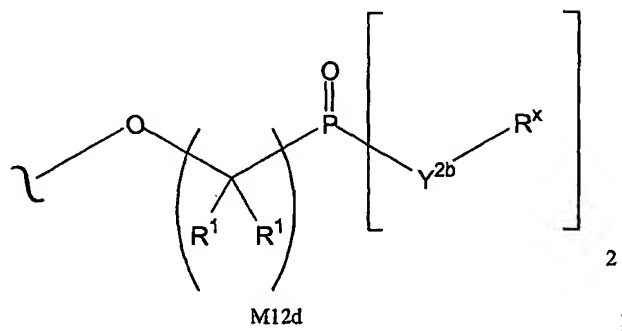
wherein Y^{1a} is O or S; and Y^{2a} is O, N(R^x) or S.

In another specific embodiment of the invention A³ is of the formula:



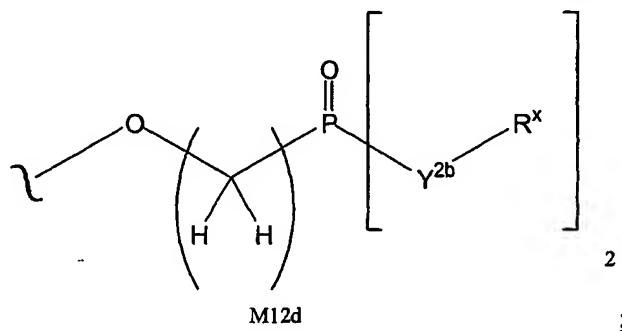
wherein Y^{2b} is O or $N(R^x)$.

In another specific embodiment of the invention A^3 is of the formula:



5 wherein Y^{2b} is O or $N(R^x)$; and M12d is 1, 2, 3, 4, 5, 6, 7 or 8.

In another specific embodiment of the invention A^3 is of the formula:

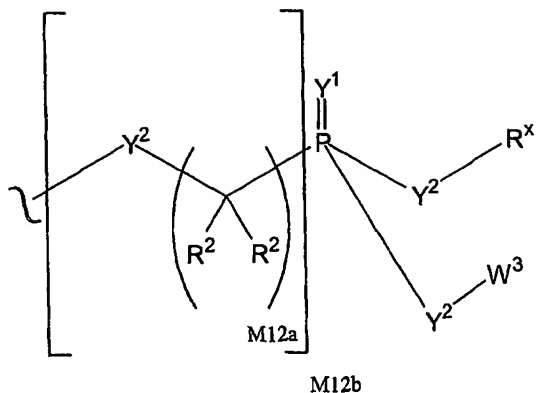


wherein Y^{2b} is O or $N(R^x)$; and M12d is 1, 2, 3, 4, 5, 6, 7 or 8.

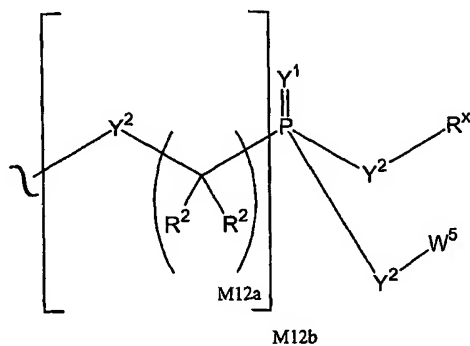
In another specific embodiment of the invention M12d is 1.

10

In another specific embodiment of the invention A^3 is of the formula:



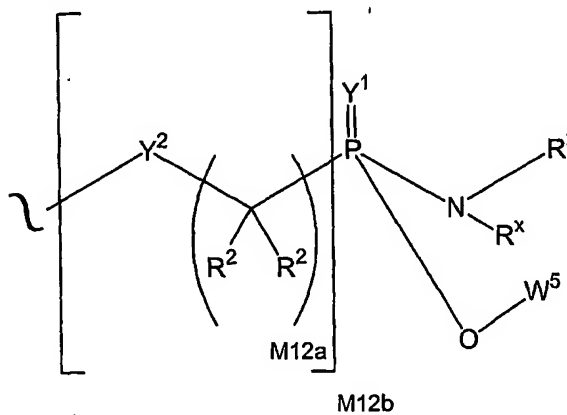
In another specific embodiment of the invention A³ is of the formula:



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In another specific embodiment of the invention W⁵ is a carbocycle.

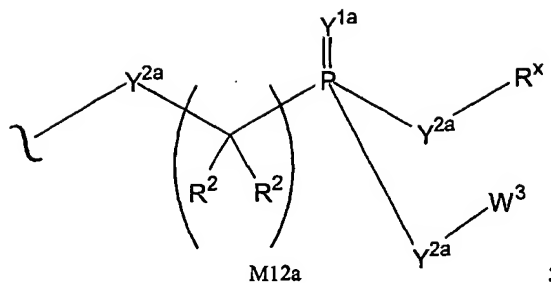
In another specific embodiment of the invention A³ is of the formula:



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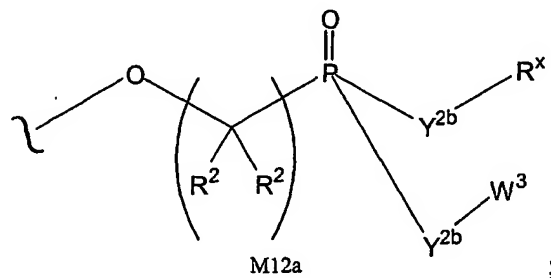
In another specific embodiment of the invention W⁵ is phenyl.

In another specific embodiment of the invention A³ is of the formula:



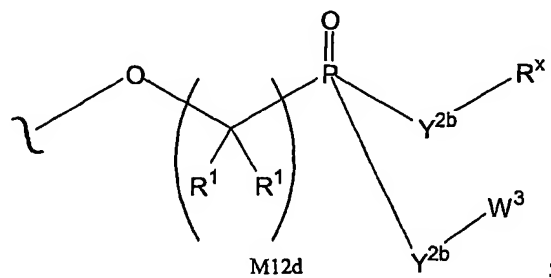
wherein Y^{1a} is O or S; and Y^{2a} is O, $N(R^x)$ or S.

In another specific embodiment of the invention A^3 is of the formula:



5 wherein Y^{2b} is O or $N(R^x)$.

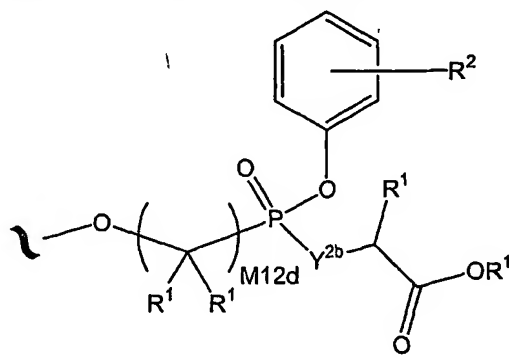
In another specific embodiment of the invention A^3 is of the formula:



wherein Y^{2b} is O or $N(R^x)$; and M12d is 1, 2, 3, 4, 5, 6, 7 or 8.

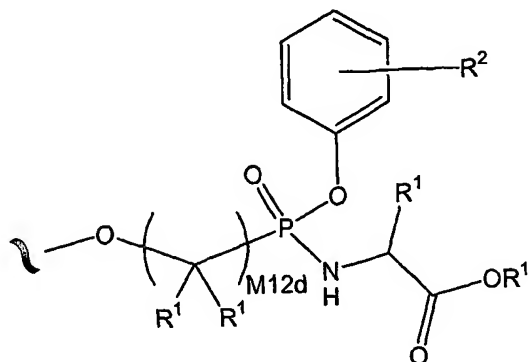
In another specific embodiment of the invention R^1 is H.

10 In another specific embodiment of the invention A^3 is of the formula:

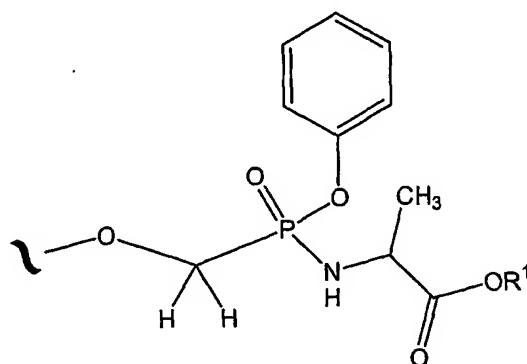


wherein the phenyl carbocycle is substituted with 0, 1, 2, or 3 R^2 groups.

In another specific embodiment of the invention A³ is of the formula:

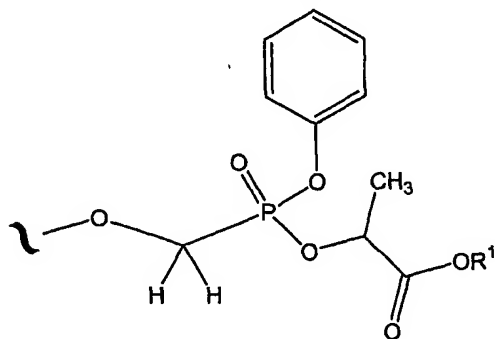


In another specific embodiment of the invention A³ is of the formula:



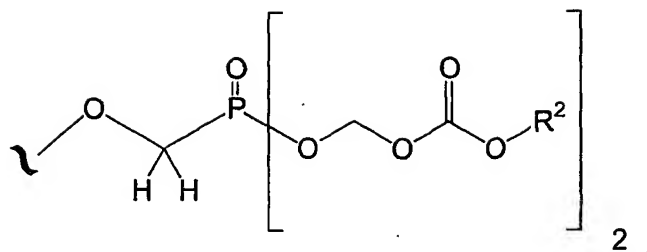
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In another specific embodiment of the invention A³ is of the formula:

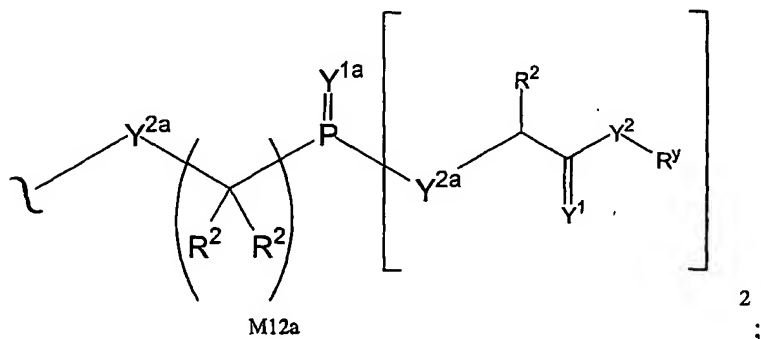


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In another specific embodiment of the invention A³ is of the formula:

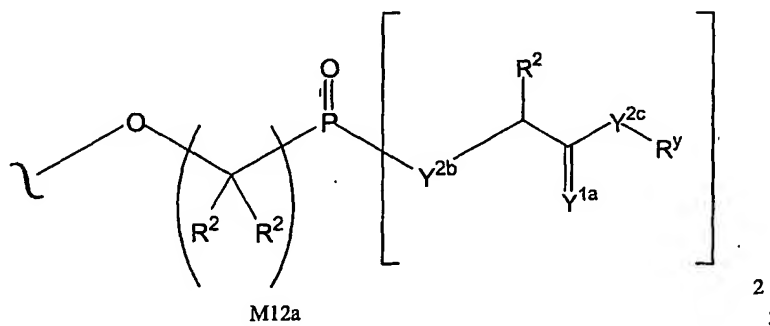


In another specific embodiment of the invention A³ is of the formula:



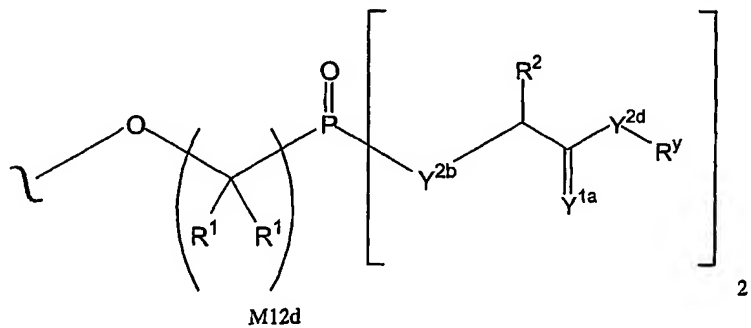
5 wherein Y^{1a} is O or S; and Y^{2a} is O, N(R²) or S.

In another specific embodiment of the invention A³ is of the formula:



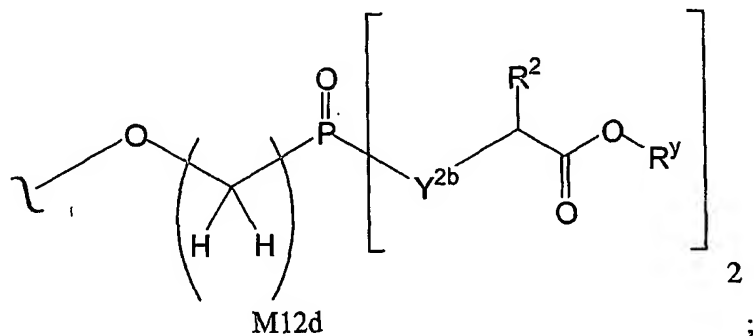
wherein Y^{1a} is O or S; Y^{2b} is O or N(R²); and Y^{2c} is O, N(R^y) or S.

10 In another specific embodiment of the invention A³ is of the formula:



wherein Y^{1a} is O or S; Y^{2b} is O or $N(R^2)$; Y^{2d} is O or $N(R^y)$; and M12d is 1, 2, 3, 4, 5, 6, 7 or 8.

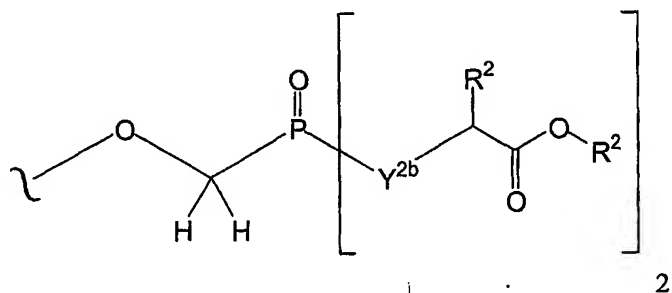
In another specific embodiment of the invention A^3 is of the formula:



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wherein Y^{2b} is O or $N(R^2)$; and M12d is 1, 2, 3, 4, 5, 6, 7 or 8.

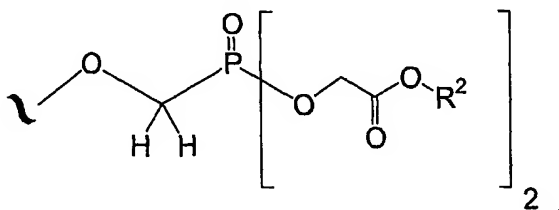
In another specific embodiment of the invention A^3 is of the formula:



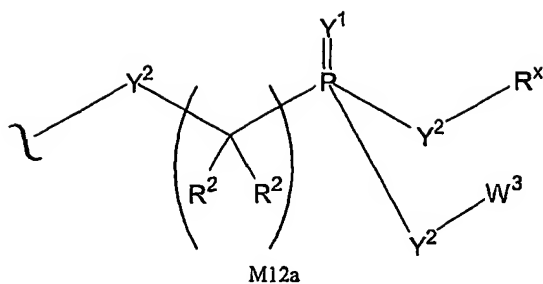
wherein Y^{2b} is O or $N(R^2)$.

10

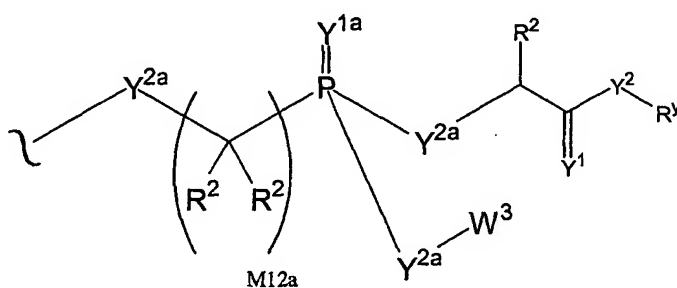
In another specific embodiment of the invention A^3 is of the formula:



In another specific embodiment of the invention A^3 is of the formula:



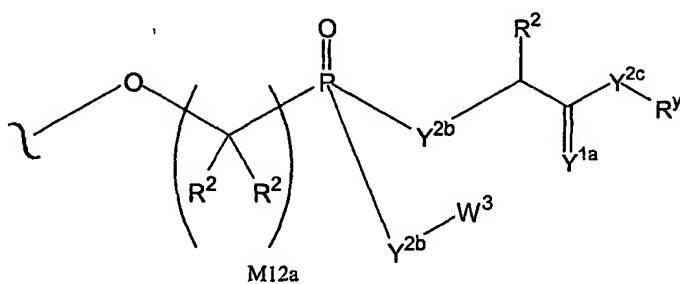
In another specific embodiment of the invention A^3 is of the formula:



5

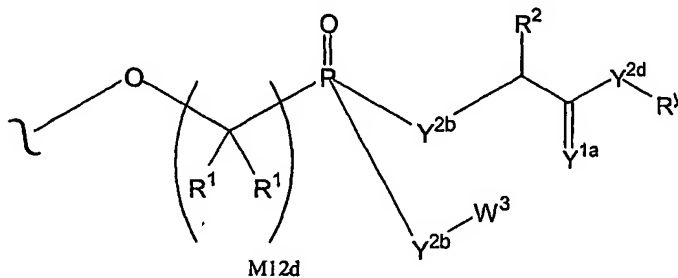
wherein Y^{1a} is O or S; and Y^{2a} is O, $N(R^2)$ or S.

In another specific embodiment of the invention A^3 is of the formula:



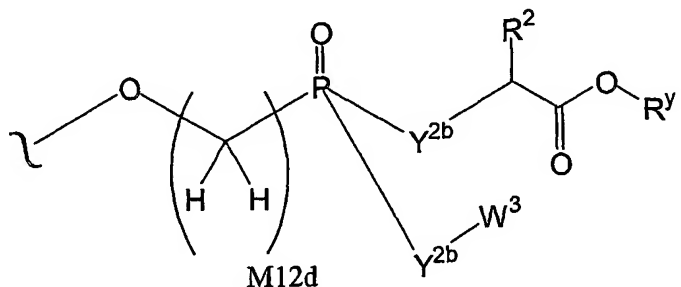
10 wherein Y^{1a} is O or S; Y^{2b} is O or $N(R^2)$; and Y^{2c} is O, $N(R^y)$ or S.

In another specific embodiment of the invention A^3 is of the formula:



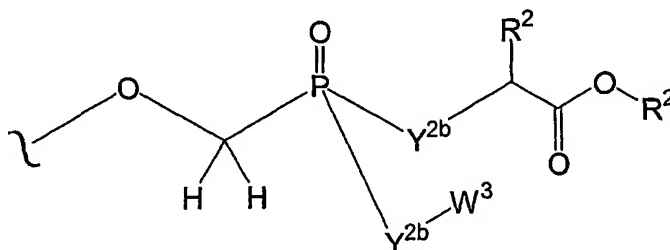
wherein Y^{1a} is O or S; Y^{2b} is O or N(R^2); Y^{2d} is O or N(R^y); and M12d is 1, 2, 3, 4, 5, 6, 7 or 8.

In another specific embodiment of the invention A^3 is of the formula:



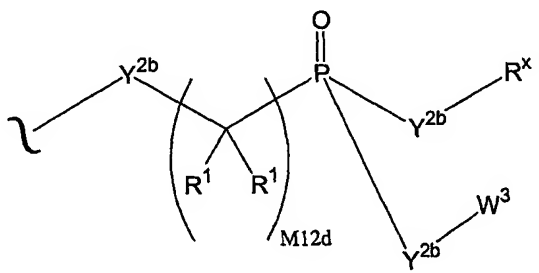
5 wherein Y^{2b} is O or N(R^2); and M12d is 1, 2, 3, 4, 5, 6, 7 or 8.

In another specific embodiment of the invention A^3 is of the formula:



wherein Y^{2b} is O or N(R^2).

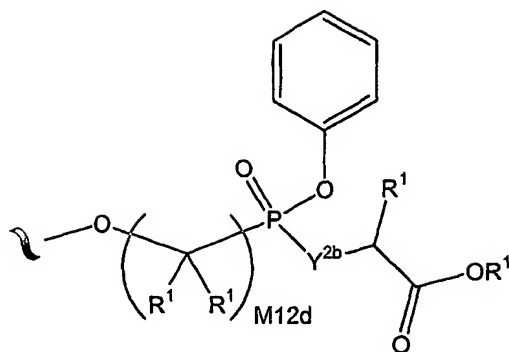
In another specific embodiment of the invention A^3 is of the formula:



10

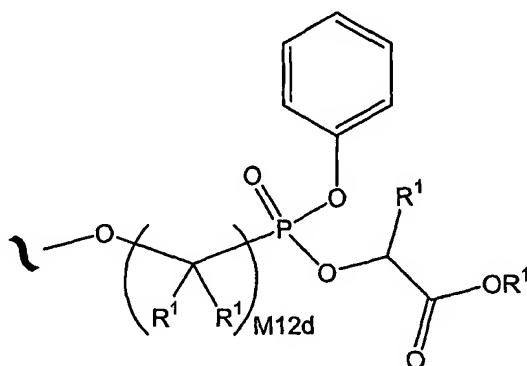
wherein: Y^{2b} is O or N(R^x); and M12d is 1, 2, 3, 4, 5, 6, 7 or 8.

In another specific embodiment of the invention A^3 is of the formula:



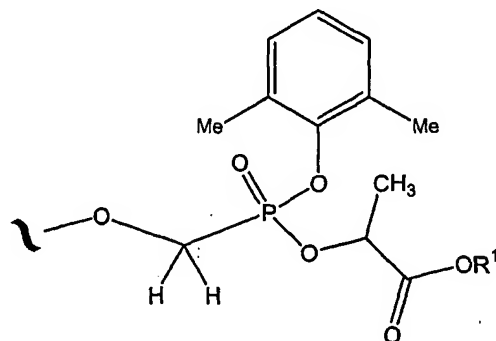
wherein the phenyl carbocycle is substituted with 0, 1, 2, or 3 R^2 groups.

In another specific embodiment of the invention A^3 is of the formula:

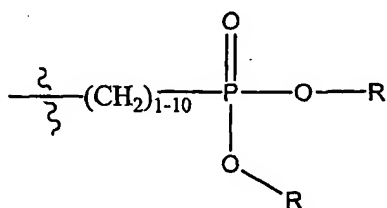


5 wherein the phenyl carbocycle is substituted with 0, 1, 2, or 3 R^2 groups.

In another specific embodiment of the invention A^3 is of the formula:

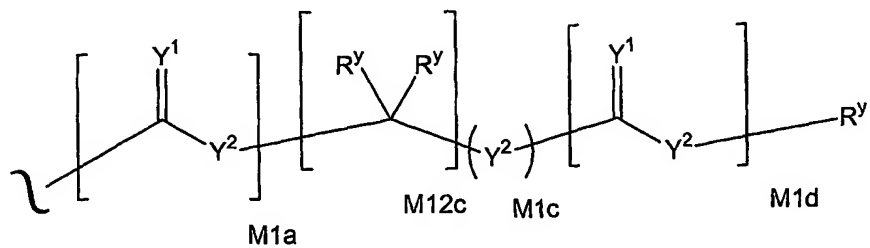


In a specific embodiment of the invention A^0 is of the formula:



wherein each R is independently (C₁-C₆)alkyl.

In a specific embodiment of the invention R^x is independently H, R¹, W³, a protecting group, or the formula:



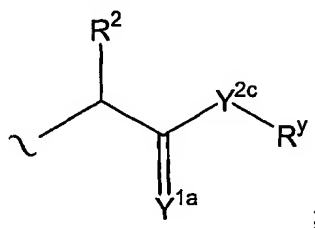
5 wherein:

R^y is independently H, W³, R² or a protecting group;

R¹ is independently H or alkyl of 1 to 18 carbon atoms;

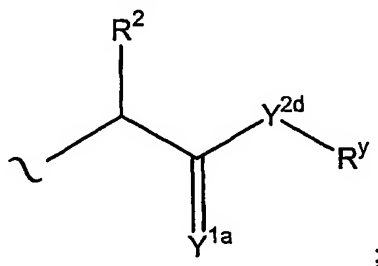
R² is independently H, R¹, R³ or R⁴ wherein each R⁴ is independently substituted with 0 to 3 R³ groups or taken together at a carbon atom, two R² groups form a ring of 3 to 8 carbons and the ring may be substituted with 0 to 3 R³ groups;

In a specific embodiment of the invention R^x is of the formula:



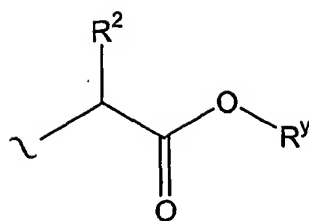
wherein Y^{1a} is O or S; and Y^{2c} is O, N(R^y) or S.

15 In a specific embodiment of the invention R^x is of the formula:



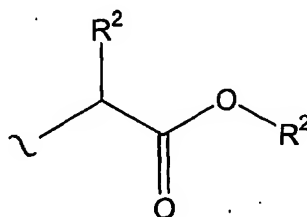
wherein Y^{1a} is O or S; and Y^{2d} is O or N(R^y).

In a specific embodiment of the invention R^x is of the formula:

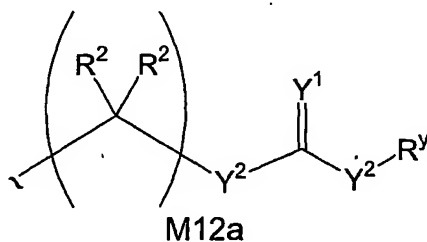


In a specific embodiment of the invention R^y is hydrogen or alkyl of 1 to 10 carbons.

5 In a specific embodiment of the invention R^x is of the formula:

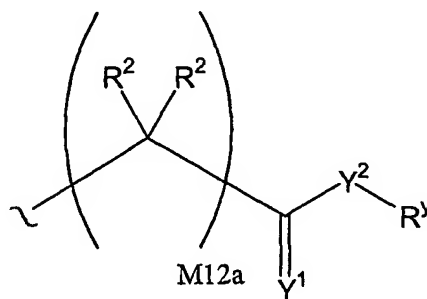


In a specific embodiment of the invention R^x is of the formula:



10

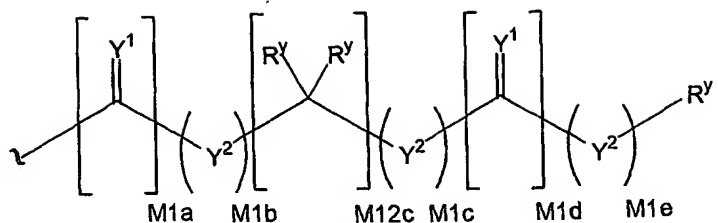
In a specific embodiment of the invention R^x is of the formula:



In a specific embodiment of the invention Y^1 is O or S.

15 In a specific embodiment of the invention Y^2 is O, $N(R^y)$ or S.

In one specific embodiment of the invention R^x is a group of the formula:



wherein:

m1a, m1b, m1c, m1d and m1e are independently 0 or 1;

5 m12c is 0, 1, 2, 3, 4, 5, 6, 7, 8, 9, 10, 11 or 12;

R^y is H, W³, R² or a protecting group;

provided that:

if m1a, m12c, and m1d are 0, then m1b, m1c and m1e are 0;

if m1a and m12c are 0 and m1d is not 0, then m1b and m1c are 0;

10 if m1a and m1d are 0 and m12c is not 0, then m1b and at least one of m1c and m1e are 0;

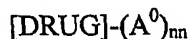
if m1a is 0 and m12c and m1d are not 0, then m1b is 0;

if m12c and m1d are 0 and m1a is not 0, then at least two of m1b, m1c and m1e are 0;

15 if m12c is 0 and m1a and m1d are not 0, then at least one of m1b and m1c are 0; and

if m1d is 0 and m1a and m12c are not 0, then at least one of m1c and m1e are 0.

20 In another specific embodiment, the invention provides a compound of the formula:



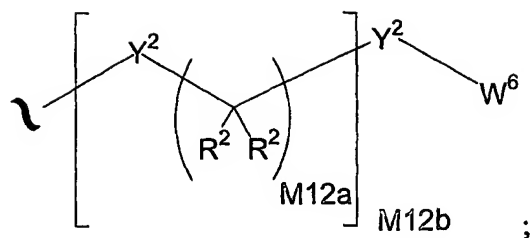
or a pharmaceutically acceptable salt thereof wherein,

25 DRUG is a compound of any one of formulae 500-547;

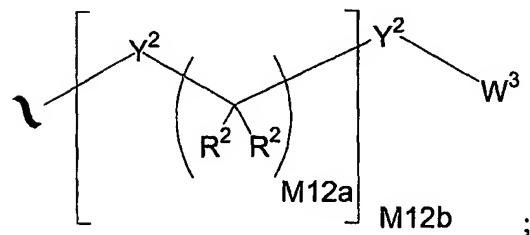
nn is 1, 2, or 3;

A⁰ is A¹, A² or W³ with the proviso that the compound includes at least one A¹;

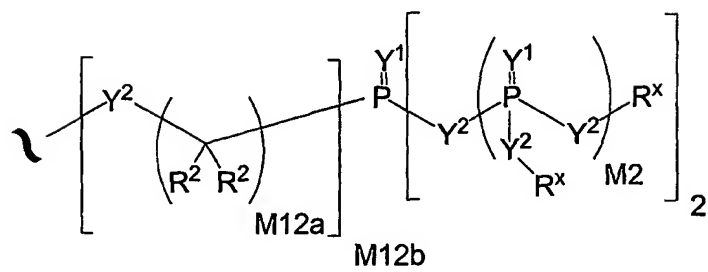
A¹ is:



A² is:



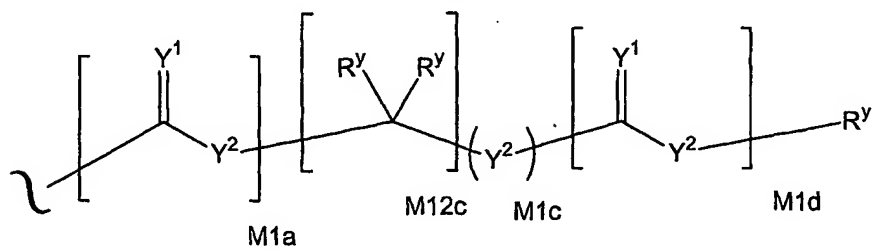
5 A³ is:



Y¹ is independently O, S, N(R^x), N(O)(R^x), N(OR^x), N(O)(OR^x), or N(N(R^x)(R^x));

Y² is independently a bond, O, N(R^x), N(O)(R^x), N(OR^x), N(O)(OR^x),
 10 N(N(R^x)(R^x)), -S(O)_{M2}-, or -S(O)_{M2}-S(O)_{M2}-;

R^x is independently H, R¹, W³, a protecting group, or the formula:



wherein:

R^y is independently H, W³, R² or a protecting group;

15 R¹ is independently H or alkyl of 1 to 18 carbon atoms;

R^2 is independently H, R^1 , R^3 or R^4 wherein each R^4 is independently substituted with 0 to 3 R^3 groups or taken together at a carbon atom, two R^2 groups form a ring of 3 to 8 carbons and the ring may be substituted with 0 to 3 R^3 groups;

5 R^3 is R^{3a} , R^{3b} , R^{3c} or R^{3d} , provided that when R^3 is bound to a heteroatom, then R^3 is R^{3c} or R^{3d} ;

R^{3a} is F, Cl, Br, I, -CN, N_3 or $-NO_2$;

R^{3b} is Y^1 ;

10 R^{3c} is $-R^x$, $-N(R^x)(R^x)$, $-SR^x$, $-S(O)R^x$, $-S(O)_2R^x$, $-S(O)(OR^x)$, $-S(O)_2(OR^x)$, $-OC(Y^1)R^x$, $-OC(Y^1)OR^x$, $-OC(Y^1)(N(R^x)(R^x))$, $-SC(Y^1)R^x$, $-SC(Y^1)OR^x$, $-SC(Y^1)(N(R^x)(R^x))$, $-N(R^x)C(Y^1)R^x$, $-N(R^x)C(Y^1)OR^x$, or $-N(R^x)C(Y^1)(N(R^x)(R^x))$;

R^{3d} is $-C(Y^1)R^x$, $-C(Y^1)OR^x$ or $-C(Y^1)(N(R^x)(R^x))$;

15 R^4 is an alkyl of 1 to 18 carbon atoms, alkenyl of 2 to 18 carbon atoms, or alkynyl of 2 to 18 carbon atoms;

R^5 is R^4 wherein each R^4 is substituted with 0 to 3 R^3 groups;

R^{5a} is independently alkylene of 1 to 18 carbon atoms, alkenylene of 2 to 18 carbon atoms, or alkynylene of 2-18 carbon atoms any one of which alkylene, alkenylene or alkynylene is substituted with 0-3 R^3 groups;

20 W^3 is W^4 or W^5 ;

W^4 is R^5 , $-C(Y^1)R^5$, $-C(Y^1)W^5$, $-SO_2R^5$, or $-SO_2W^5$;

W^5 is carbocycle or heterocycle wherein W^5 is independently substituted with 0 to 3 R^2 groups;

W^6 is W^3 independently substituted with 1, 2, or 3 A^3 groups;

25 $M2$ is 0, 1 or 2;

$M12a$ is 1, 2, 3, 4, 5, 6, 7, 8, 9, 10, 11 or 12;

$M12b$ is 0, 1, 2, 3, 4, 5, 6, 7, 8, 9, 10, 11 or 12;

$M1a$, $M1c$, and $M1d$ are independently 0 or 1; and

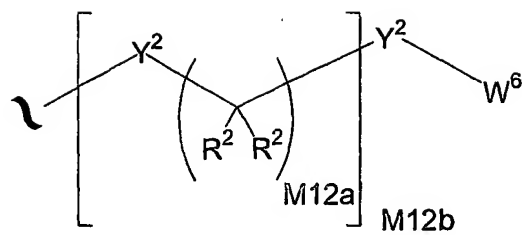
$M12c$ is 0, 1, 2, 3, 4, 5, 6, 7, 8, 9, 10, 11 or 12.

30

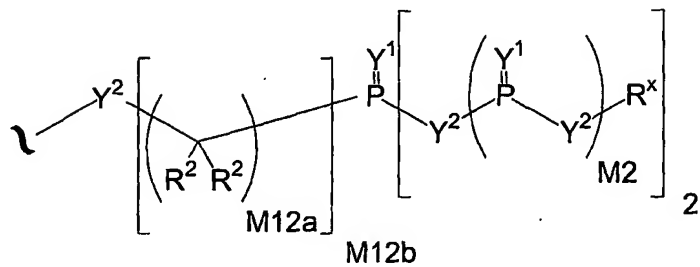
In another embodiment, the invention provides a compound of any one of formulae 1-151 wherein:

A^0 is A^1 ;

A¹ is:



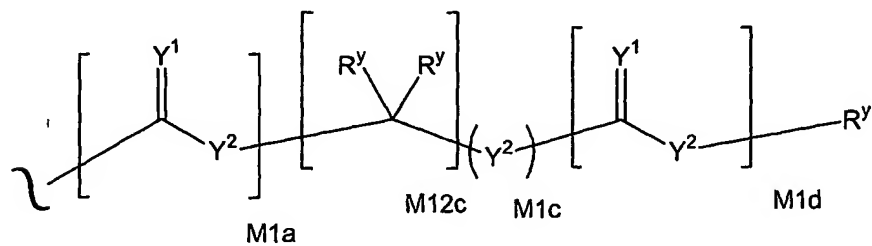
A³ is:



5 Y¹ is independently O, S, N(R^x), N(O)(R^x), N(OR^x), N(O)(OR^x), or N(N(R^x)(R^x));

Y² is independently a bond, O, N(R^x), N(O)(R^x), N(OR^x), N(O)(OR^x), N(N(R^x)(R^x)), -S(O)_{M2}-, or -S(O)_{M2}-S(O)_{M2}-;

R^x is independently H, W³, a protecting group, or the formula:



10

R^y is independently H, W³, R² or a protecting group;

R¹ is independently H or alkyl of 1 to 18 carbon atoms;

R² is independently H, R³ or R⁴ wherein each R⁴ is independently substituted with 0 to 3 R³ groups;

15 R³ is R^{3a}, R^{3b}, R^{3c} or R^{3d}, provided that when R³ is bound to a heteroatom, then R³ is R^{3c} or R^{3d};

R^{3a} is F, Cl, Br, I, -CN, N₃ or -NO₂;

R^{3b} is Y¹;

20 R^{3c} is -R^x, -N(R^x)(R^x), -SR^x, -S(O)R^x, -S(O)₂R^x, -S(O)(OR^x), -S(O)₂(OR^x), -OC(Y¹)R^x, -OC(Y¹)OR^x, -OC(Y¹)(N(R^x)(R^x)), -SC(Y¹)R^x, -

SC(Y¹)OR^x, -SC(Y¹)(N(R^x)(R^x)), -N(R^x)C(Y¹)R^x, -N(R^x)C(Y¹)OR^x, or -N(R^x)C(Y¹)(N(R^x)(R^x));

R^{3d} is -C(Y¹)R^x, -C(Y¹)OR^x or -C(Y¹)(N(R^x)(R^x));

R⁴ is an alkyl of 1 to 18 carbon atoms, alkenyl of 2 to 18 carbon atoms,

5 or alkynyl of 2 to 18 carbon atoms;

R⁵ is R⁴ wherein each R⁴ is substituted with 0 to 3 R³ groups;

R^{5a} is independently alkylene of 1 to 18 carbon atoms, alkenylene of 2 to 18 carbon atoms, or alkynylene of 2-18 carbon atoms any one of which alkylene, alkenylene or alkynylene is substituted with 0-3 R³ groups;

10 W³ is W⁴ or W⁵;

W⁴ is R⁵, -C(Y¹)R⁵, -C(Y¹)W⁵, -SO₂R⁵, or -SO₂W⁵;

W⁵ is carbocycle or heterocycle wherein W⁵ is independently substituted with 0 to 3 R² groups;

W⁶ is W³ independently substituted with 1, 2, or 3 A³ groups;

15 M2 is 0, 1 or 2;

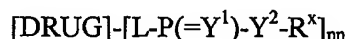
M12a is 1, 2, 3, 4, 5, 6, 7, 8, 9, 10, 11 or 12;

M12b is 0, 1, 2, 3, 4, 5, 6, 7, 8, 9, 10, 11 or 12;

M1a, M1c, and M1d are independently 0 or 1; and

M12c is 0, 1, 2, 3, 4, 5, 6, 7, 8, 9, 10, 11 or 12.

20 In another specific embodiment, the invention provides a compound of the formula:



25 or a pharmaceutically acceptable salt thereof wherein,

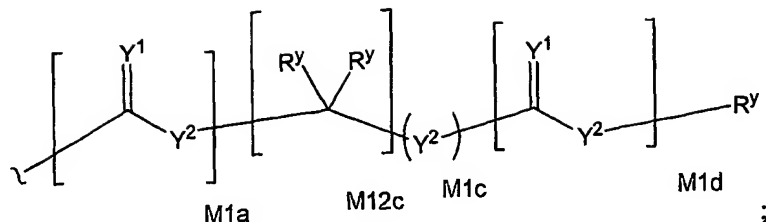
DRUG is a compound of any one of formulae 500-547;

Y¹ is independently O, S, N(R^x), N(O)(R^x), N(OR^x), N(O)(OR^x), or N(N(R^x)(R^x));

Y² is independently a bond, O, N(R^x), N(O)(R^x), N(OR^x), N(O)(OR^x),

30 N(N(R^x)(R^x)), -S(O)_{M2}-, or -S(O)_{M2}-S(O)_{M2}-;

R^x is independently H, W³, a protecting group, or the formula:



R^y is independently H, W^3 , R^2 or a protecting group;

R^2 is independently H, R^3 or R^4 wherein each R^4 is independently substituted with 0 to 3 R^3 groups;

5 R^3 is R^{3a} , R^{3b} , R^{3c} or R^{3d} , provided that when R^3 is bound to a heteroatom, then R^3 is R^{3c} or R^{3d} ;

R^{3a} is F, Cl, Br, I, -CN, N_3 or $-NO_2$;

R^{3b} is Y^1 ;

R^{3c} is $-R^x$, $-N(R^x)(R^x)$, $-SR^x$, $-S(O)R^x$, $-S(O)_2R^x$, $-S(O)(OR^x)$, -

10 $S(O)_2(OR^x)$,

$-OC(Y^1)R^x$, $-OC(Y^1)OR^x$, $-OC(Y^1)(N(R^x)(R^x))$, $-SC(Y^1)R^x$, $-SC(Y^1)OR^x$,

$-SC(Y^1)(N(R^x)(R^x))$, $-N(R^x)C(Y^1)R^x$, $-N(R^x)C(Y^1)OR^x$, or -

$N(R^x)C(Y^1)(N(R^x)(R^x))$;

R^{3d} is $-C(Y^1)R^x$, $-C(Y^1)OR^x$ or $-C(Y^1)(N(R^x)(R^x))$;

15 R^4 is an alkyl of 1 to 18 carbon atoms, alkenyl of 2 to 18 carbon atoms, or alkynyl of 2 to 18 carbon atoms;

R^5 is R^4 wherein each R^4 is substituted with 0 to 3 R^3 groups;

W^3 is W^4 or W^5 ;

W^4 is R^5 , $-C(Y^1)R^5$, $-C(Y^1)W^5$, $-SO_2R^5$, or $-SO_2W^5$;

20 W^5 is carbocycle or heterocycle wherein W^5 is independently substituted with 0 to 3 R^2 groups;

M2 is 1, 2, or 3;

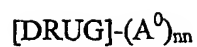
M1a, M1c, and M1d are independently 0 or 1;

M12c is 0, 1, 2, 3, 4, 5, 6, 7, 8, 9, 10, 11 or 12;

25 n is 1, 2, or 3; and

L is a linking group.

In another specific embodiment, the invention provides a compound of which is a compound of the formula:



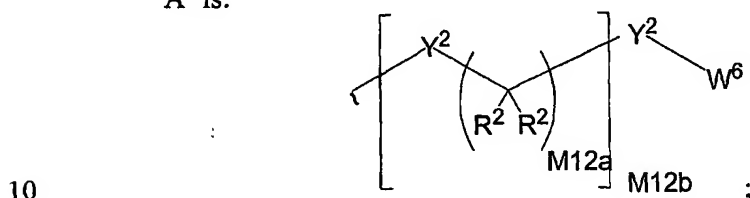
or a pharmaceutically acceptable salt thereof wherein,

DRUG is a compound of any one of formulae 500-547;

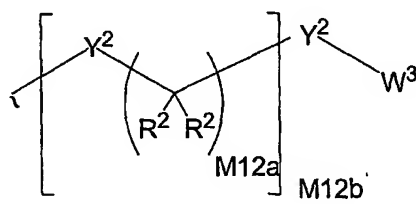
5 mn is 1, 2, or 3;

A^0 is A^1 , A^2 , or W^3 with the proviso that the compound includes at least one A^1 ;

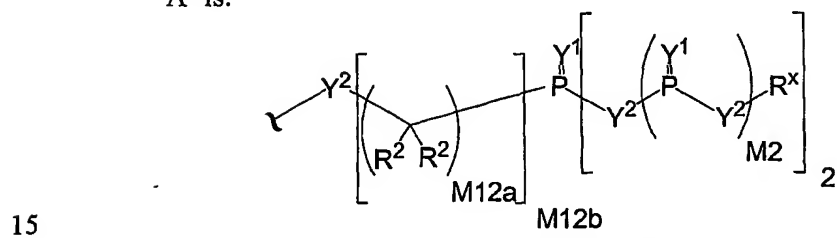
A^1 is:



A^2 is:



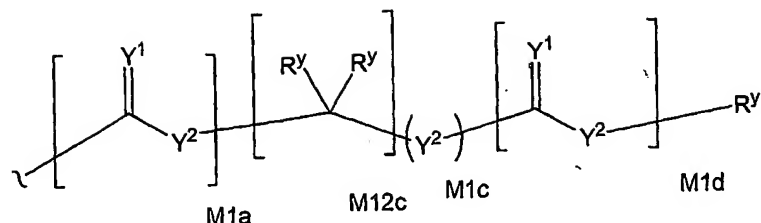
A^3 is:



Y^1 is independently O, S, $\text{N}(\text{R}^x)$, $\text{N}(\text{O})(\text{R}^x)$, $\text{N}(\text{OR}^x)$, $\text{N}(\text{O})(\text{OR}^x)$, or $\text{N}(\text{N}(\text{R}^x)(\text{R}^x))$;

Y^2 is independently a bond, O, $\text{N}(\text{R}^x)$, $\text{N}(\text{O})(\text{R}^x)$, $\text{N}(\text{OR}^x)$, $\text{N}(\text{O})(\text{OR}^x)$, $\text{N}(\text{N}(\text{R}^x)(\text{R}^x))$, $-\text{S}(\text{O})_{\text{M}2-}$, or $-\text{S}(\text{O})_{\text{M}2-}\text{S}(\text{O})_{\text{M}2-}$;

20 R^x is independently H, W^3 , a protecting group, or the formula:



R^Y is independently H, W^3 , R^2 or a protecting group;

R^2 is independently H, R^3 or R^4 wherein each R^4 is independently substituted with 0 to 3 R^3 groups;

- 5 R^3 is R^{3a} , R^{3b} , R^{3c} or R^{3d} , provided that when R^3 is bound to a heteroatom, then R^3 is R^{3c} or R^{3d} ;

R^{3a} is F, Cl, Br, I, -CN, N_3 or -NO₂;

R^{3b} is Y^1 ;

R^{3c} is - R^x , - $N(R^x)(R^x)$, - SR^x , - $S(O)R^x$, - $S(O)_2R^x$, - $S(O)(OR^x)$, -

- 10 $S(O)_2(OR^x)$,

- $OC(Y^1)R^x$, - $OC(Y^1)OR^x$, - $OC(Y^1)(N(R^x)(R^x))$, - $SC(Y^1)R^x$, - $SC(Y^1)OR^x$,

- $SC(Y^1)(N(R^x)(R^x))$, - $N(R^x)C(Y^1)R^x$, - $N(R^x)C(Y^1)OR^x$, or -

$N(R^x)C(Y^1)(N(R^x)(R^x))$;

R^{3d} is - $C(Y^1)R^x$, - $C(Y^1)OR^x$ or - $C(Y^1)(N(R^x)(R^x))$;

- 15 R^4 is an alkyl of 1 to 18 carbon atoms, alkenyl of 2 to 18 carbon atoms, or alkynyl of 2 to 18 carbon atoms;

R^5 is R^4 wherein each R^4 is substituted with 0 to 3 R^3 groups;

W^3 is W^4 or W^5 ;

W^4 is R^5 , - $C(Y^1)R^5$, - $C(Y^1)W^5$, -SO₂ R^5 , or -SO₂ W^5 ;

- 20 W^5 is carbocycle or heterocycle wherein W^5 is independently substituted with 0 to 3 R^2 groups;

W^6 is W^3 independently substituted with 1, 2, or 3 A^3 groups;

M2 is 0, 1 or 2;

M12a is 1, 2, 3, 4, 5, 6, 7, 8, 9, 10, 11 or 12;

- 25 M12b is 0, 1, 2, 3, 4, 5, 6, 7, 8, 9, 10, 11 or 12;

M1a, M1c, and M1d are independently 0 or 1; and

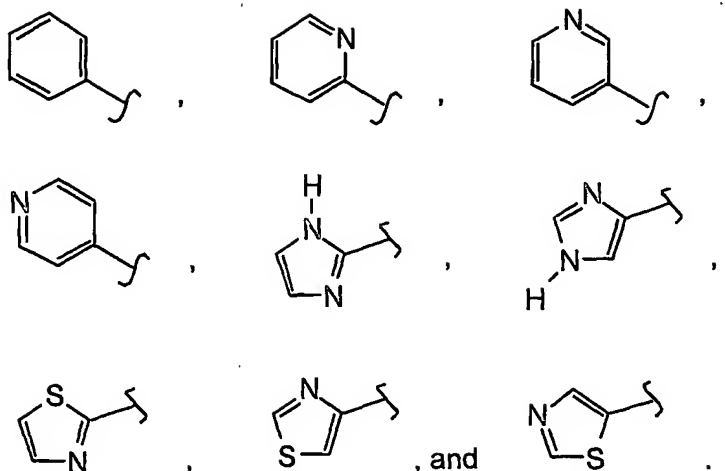
M12c is 0, 1, 2, 3, 4, 5, 6, 7, 8, 9, 10, 11 or 12.

In compounds of the invention W^5 carbocycles and W^5 heterocycles may

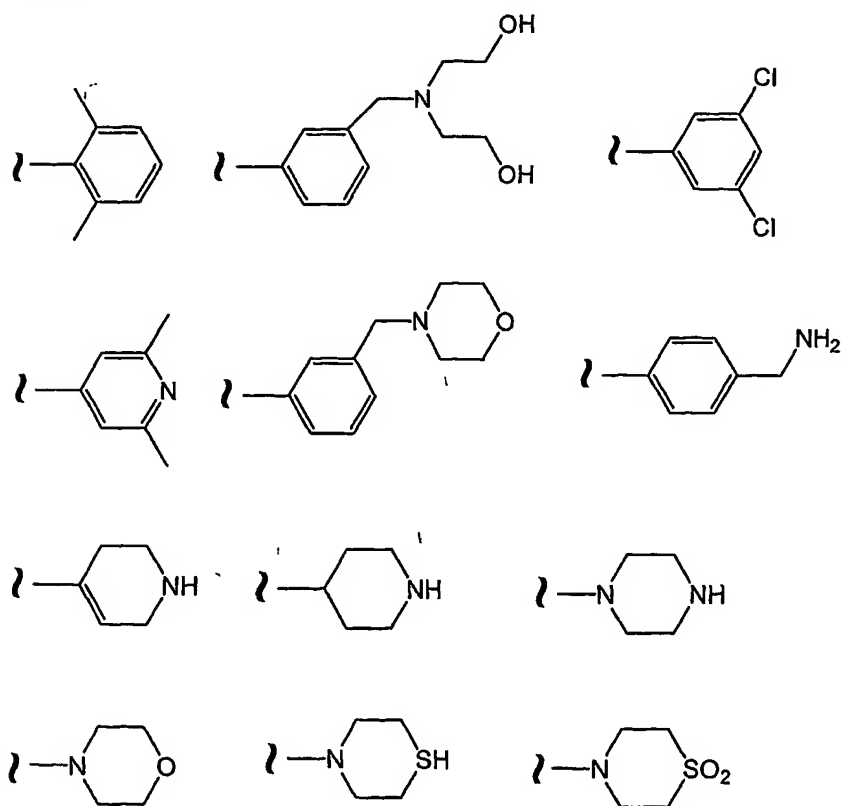
be independently substituted with 0 to 3 R^2 groups. W^5 may be a saturated, unsaturated or aromatic ring comprising a mono- or bicyclic carbocycle or heterocycle. W^5 may have 3 to 10 ring atoms, e.g., 3 to 7 ring atoms. The W^5 rings are saturated when containing 3 ring atoms, saturated or mono-unsaturated when containing 4 ring atoms, saturated, or mono- or di-unsaturated when containing 5 ring atoms, and saturated, mono- or di-unsaturated, or aromatic when containing 6 ring atoms.

A W^5 heterocycle may be a monocycle having 3 to 7 ring members (2 to 6 carbon atoms and 1 to 3 heteroatoms selected from N, O, P, and S) or a bicycle having 7 to 10 ring members (4 to 9 carbon atoms and 1 to 3 heteroatoms selected from N, O, P, and S). W^5 heterocyclic monocycles may have 3 to 6 ring atoms (2 to 5 carbon atoms and 1 to 2 heteroatoms selected from N, O, and S); or 5 or 6 ring atoms (3 to 5 carbon atoms and 1 to 2 heteroatoms selected from N and S). W^5 heterocyclic bicycles have 7 to 10 ring atoms (6 to 9 carbon atoms and 1 to 2 heteroatoms selected from N, O, and S) arranged as a bicyclo [4,5], [5,5], [5,6], or [6,6] system; or 9 to 10 ring atoms (8 to 9 carbon atoms and 1 to 2 hetero atoms selected from N and S) arranged as a bicyclo [5,6] or [6,6] system. The W^5 heterocycle may be bonded to Y^2 through a carbon, nitrogen, sulfur or other atom by a stable covalent bond.

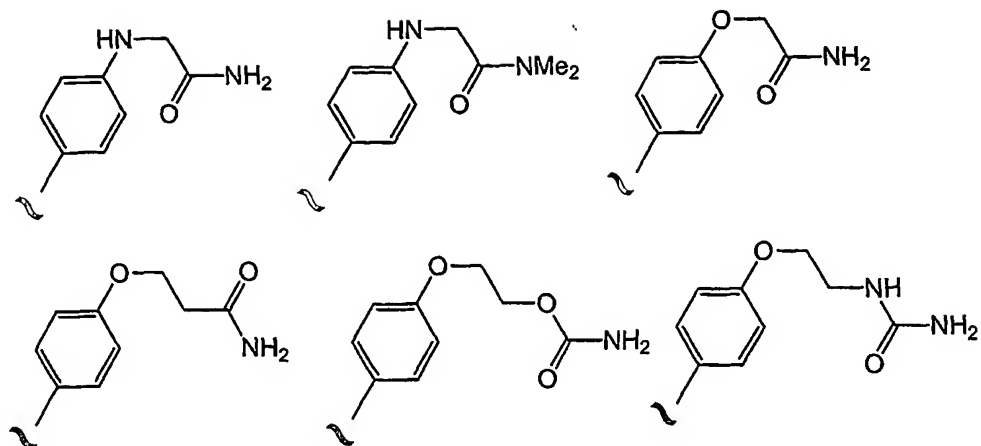
W^5 heterocycles include for example, pyridyl, dihydropyridyl isomers, piperidine, pyridazinyl, pyrimidinyl, pyrazinyl, s-triazinyl, oxazolyl, imidazolyl, thiazolyl, isoxazolyl, pyrazolyl, isothiazolyl, furanyl, thiofuranyl, thienyl, and pyrrolyl. W^5 also includes, but is not limited to, examples such as:



W⁵ carbocycles and heterocycles may be independently substituted with 0 to 3 R² groups, as defined above. For example, substituted W⁵ carbocycles include:



Examples of substituted phenyl carbocycles include:



Linking Groups and Linkers

The invention provides conjugates that comprise an immuno-modulatory compound that is linked to one or more phosphonate groups either directly (e.g. through a covalent bond) or through a linking group (i.e. a linker). The nature of the linker is not critical provided it does not interfere with the ability of the phosphonate containing compound to function as a therapeutic agent. The phosphonate or the linker can be linked to the compound (e.g. a compound of Formulae 500-547) at any synthetically feasible position on the compound by removing a hydrogen or any portion of the compound to provide an open valence for attachment of the phosphonate or the linker.

In one embodiment of the invention the linking group or linker (which can be designated "L") can include all or a portions of the group A^0 , A^1 , A^2 , or W^3 described herein.

In another embodiment of the invention the linking group or linker has a molecular weight of from about 20 daltons to about 400 daltons.

In another embodiment of the invention the linking group or linker has a length of about 5 angstroms to about 300 angstroms.

In another embodiment of the invention the linking group or linker separates the DRUG and a $\text{P}(=\text{Y}^1)$ residue by about 5 angstroms to about 200 angstroms, inclusive, in length.

In another embodiment of the invention the linking group or linker is a divalent, branched or unbranched, saturated or unsaturated, hydrocarbon chain, having from 2 to 25 carbon atoms, wherein one or more (e.g. 1, 2, 3, or 4) of the

carbon atoms is optionally replaced by (-O-), and wherein the chain is optionally substituted on carbon with one or more (e.g. 1, 2, 3, or 4) substituents selected from (C₁-C₆)alkoxy, (C₃-C₆)cycloalkyl, (C₁-C₆)alkanoyl, (C₁-C₆)alkanoyloxy, (C₁-C₆)alkoxycarbonyl, (C₁-C₆)alkylthio, azido, cyano, nitro, halo, hydroxy, 5 oxo (=O), carboxy, aryl, aryloxy, heteroaryl, and heteroaryloxy.

In another embodiment of the invention the linking group or linker is of the formula W-A wherein A is (C₁-C₂₄)alkyl, (C₂-C₂₄)alkenyl, (C₂-C₂₄)alkynyl, (C₃-C₈)cycloalkyl, (C₆-C₁₀)aryl or a combination thereof, wherein W is - N(R)C(=O)-, -C(=O)N(R)-, -OC(=O)-, -C(=O)O-, -O-, -S-, -S(O)-, -S(O)₂-, - 10 N(R)-, -C(=O)-, or a direct bond; wherein each R is independently H or (C₁-C₆)alkyl.

In another embodiment of the invention the linking group or linker is a divalent radical formed from a peptide.

In another embodiment of the invention the linking group or linker is a 15 divalent radical formed from an amino acid.

In another embodiment of the invention the linking group or linker is a divalent radical formed from poly-L-glutamic acid, poly-L-aspartic acid, poly-L-histidine, poly-L-ornithine, poly-L-serine, poly-L-threonine, poly-L-tyrosine, poly-L-leucine, poly-L-lysine-L-phenylalanine, poly-L-lysine or poly-L-lysine- 20 L-tyrosine.

In another embodiment of the invention the linking group or linker is of the formula W-(CH₂)_n wherein, n is between about 1 and about 10; and W is - N(R)C(=O)-, -C(=O)N(R)-, -OC(=O)-, -C(=O)O-, -O-, -S-, -S(O)-, -S(O)₂-, - C(=O)-, -N(R)-, or a direct bond; wherein each R is independently H or (C₁- 25 C₆)alkyl.

In another embodiment of the invention the linking group or linker is methylene, ethylene, or propylene.

In another embodiment of the invention the linking group or linker is attached to the phosphonate group through a carbon atom of the linker. 30

Compounds

The compounds of the invention include those with immuno-modulatory activity. The compounds of the inventions bear one or more (e.g. 1, 2, 3, or 4)

phosphonate groups, which may be a prodrug moiety.

Typically, compounds of the invention have a molecular weight of from about 400 amu to about 10,000 amu; in a specific embodiment of the invention, compounds have a molecular weight of less than about 5000 amu; in another
5 specific embodiment of the invention, compounds have a molecular weight of less than about 2500 amu; in another specific embodiment of the invention, compounds have a molecular weight of less than about 1000 amu; in another specific embodiment of the invention, compounds have a molecular weight of less than about 800 amu; in another specific embodiment of the invention,
10 compounds have a molecular weight of less than about 600 amu; and in another specific embodiment of the invention, compounds have a molecular weight of less than about 600 amu and a molecular weight of greater than about 400 amu.

The compounds of the invention also typically have a logD(polarity) less than about 5. In one embodiment the invention provides compounds having a
15 logD less than about 4; in another one embodiment the invention provides compounds having a logD less than about 3; in another one embodiment the invention provides compounds having a logD greater than about -5; in another one embodiment the invention provides compounds having a logD greater than about -3; and in another one embodiment the invention provides compounds
20 having a logD greater than about 0 and less than about 3.

Selected substituents within the compounds of the invention are present to a recursive degree. In this context, "recursive substituent" means that a substituent may recite another instance of itself. Because of the recursive nature of such substituents, theoretically, a large number may be present in any given
25 claim. For example, R^x contains a R^y substituent. R^y can be R^2 , which in turn can be R^3 . If R^3 is selected to be R^{3c} , then a second instance of R^x can be selected. One of ordinary skill in the art of medicinal chemistry understands that the total number of such substituents is reasonably limited by the desired properties of the compound intended. Such properties include, by of example
30 and not limitation, physical properties such as molecular weight, solubility or log P, application properties such as activity against the intended target, and practical properties such as ease of synthesis.

By way of example and not limitation, W^3 , R^y and R^3 are all recursive substituents in certain claims. Typically, each of these may independently occur 20, 19, 18, 17, 16, 15, 14, 13, 12, 11, 10, 9, 8, 7, 6, 5, 4, 3, 2, 1, or 0, times in a given claim. More typically, each of these may independently occur 12 or fewer
5 times in a given claim. More typically yet, W^3 will occur 0 to 8 times, R^y will occur 0 to 6 times and R^3 will occur 0 to 10 times in a given claim. Even more typically, W^3 will occur 0 to 6 times, R^y will occur 0 to 4 times and R^3 will occur 0 to 8 times in a given claim.

Recursive substituents are an intended aspect of the invention. One of
10 ordinary skill in the art of medicinal chemistry understands the versatility of such substituents. To the degree that recursive substituents are present in an claim of the invention, the total number will be determined as set forth above.

Whenever a compound described herein is substituted with more than one of the same designated group, e.g., " R^1 " or " R^{6a} ", then it will be understood
15 that the groups may be the same or different, i.e., each group is independently selected. Wavy lines indicate the site of covalent bond attachments to the adjoining groups, moieties, or atoms.

The term immuno-modulatory compound also includes pimecrolimus, everolimus, sirolimus, tacrolimus, prednisolone, VX-148, merimepodib,
20 brequinar, thalidomide, BCX-1777, revimid, diprolene, aclometasone dipropionate, hydrocortisone, dexamethasone, leflunomide, methylprednisolone suleptanate, prednisone, clobetasol, MNA-715 (FK778), SMP-114, teriflunomide, halobetasol, ciclesonide, deflazacort, medroxyprogesterone, budesonide, rimexolone, triamcinolone acetonide, fluticasone, mometasone
25 furoate, methylprednisolone aceponate, cyclosporin A, tacrolimus, mycophenolate, ANA-245, immunosuppressive macrolide, methotrexate, PNP-405, MDL-74428, 9-(3,3-dimethyl-5-phosphonopentyl) guanine, DADMe-IMMG, CP-690,550, mycophenate, cyclosporin, and mizoribine.

In one embodiment of the invention, the compound is in isolated and
30 purified form. Generally, the term "isolated and purified" means that the compound is significantly free of biological materials (e.g. blood, cells, etc.). In one specific embodiment of the invention, the term means that the compound or conjugate of the invention is at least about 50% pure by weight in a mixture; in

another specific embodiment, the term means that the compound or conjugate of the invention is at least about 75% pure by weight in a mixture; in another specific embodiment, the term means that the compound or conjugate of the invention is at least about 90% pure by weight in a mixture; in another specific embodiment, the term means that the compound or conjugate of the invention is at least about 98% pure by weight in a mixture; and in another embodiment, the term means that the compound or conjugate of the invention is at least about 99% pure by weight in a mixture. In another specific embodiment, the invention provides a compound or conjugate of the invention that has been synthetically prepared (e.g. prepared *ex vivo*).

Intracellular Targeting

The phosphonate group of the compounds of the invention may cleave *in vivo* in stages after they have reached the desired site of action, i.e. inside a cell. One mechanism of action inside a cell may entail a first cleavage, e.g. by esterase, to provide a negatively-charged "locked-in" intermediate. Cleavage of a terminal ester grouping in a compound of the invention thus affords an unstable intermediate which releases a negatively charged "locked in" intermediate.

After passage inside a cell, intracellular enzymatic cleavage or modification of the phosphonate or prodrug compound may result in an intracellular accumulation of the cleaved or modified compound by a "trapping" mechanism. The cleaved or modified compound may then be "locked-in" the cell by a significant change in charge, polarity, or other physical property change which decreases the rate at which the cleaved or modified compound can exit the cell, relative to the rate at which it entered as the phosphonate prodrug. Other mechanisms by which a therapeutic effect are achieved may be operative as well. Enzymes which are capable of an enzymatic activation mechanism with the phosphonate prodrug compounds of the invention include, but are not limited to, amidases, esterases, microbial enzymes, phospholipases, cholinesterases, and phosphatases.

From the foregoing, it will be apparent that many different drugs can be derivatized in accord with the present invention. Numerous such drugs are specifically mentioned herein. However, it should be understood that the

discussion of drug families and their specific members for derivatization according to this invention is not intended to be exhaustive, but merely illustrative.

In one embodiment of the invention, the compound is not an antiviral agent compound. In another embodiment the compound is not a nucleoside compound. In another embodiment the compound is not an IMPDH inhibitor compound. In another embodiment the compound is not an antimetabolite compound. In another embodiment the compound is not a PNP inhibitor. In another embodiment the compound is not a substituted compound of any one of formulae 500-533, 535-541, or 543-547. In another embodiment the compound is not a substituted compound of any one of one of formulae 1-104, 107-124, or 128-151.

Stereoisomers

The compounds of the invention may have chiral centers, e.g., chiral carbon or phosphorus atoms. The compounds of the invention thus include racemic mixtures of all stereoisomers, including enantiomers, diastereomers, and atropisomers. In addition, the compounds of the invention include enriched or resolved optical isomers at any or all asymmetric, chiral atoms. In other words, the chiral centers apparent from the depictions are provided as the chiral isomers or racemic mixtures. Both racemic and diastereomeric mixtures, as well as the individual optical isomers isolated or synthesized, substantially free of their enantiomeric or diastereomeric partners, are all within the scope of the invention. The racemic mixtures are separated into their individual, substantially optically pure isomers through well-known techniques such as, for example, the separation of diastereomeric salts formed with optically active adjuncts, e.g., acids or bases followed by conversion back to the optically active substances. In most instances, the desired optical isomer is synthesized by means of stereospecific reactions, beginning with the appropriate stereoisomer of the desired starting material.

The compounds of the invention can also exist as tautomeric isomers in certain cases. All though only one delocalized resonance structure may be depicted, all such forms are contemplated within the scope of the invention. For

example, ene-amine tautomers can exist for purine, pyrimidine, imidazole, guanidine, amidine, and tetrazole systems and all their possible tautomeric forms are within the scope of the invention.

Salts and Hydrates

5 The compositions of this invention optionally comprise salts of the compounds herein, especially pharmaceutically acceptable non-toxic salts containing, for example, Na^+ , Li^+ , K^+ , Ca^{+2} and Mg^{+2} . Such salts may include those derived by combination of appropriate cations such as alkali and alkaline earth metal ions or ammonium and quaternary amino ions with an acid anion
10 moiety, typically a carboxylic acid. Monovalent salts are preferred if a water soluble salt is desired.

 Metal salts typically are prepared by reacting the metal hydroxide with a compound of this invention. Examples of metal salts which are prepared in this way are salts containing Li^+ , Na^+ , and K^+ . A less soluble metal salt can be
15 precipitated from the solution of a more soluble salt by addition of the suitable metal compound.

 In addition, salts may be formed from acid addition of certain organic and inorganic acids, e.g., HCl , HBr , H_2SO_4 , H_3PO_4 or organic sulfonic acids, to basic centers, typically amines, or to acidic groups. Finally, it is to be
20 understood that the compositions herein comprise compounds of the invention in their un-ionized, as well as zwitterionic form, and combinations with stoichiometric amounts of water as in hydrates.

 Also included within the scope of this invention are the salts of the parental compounds with one or more amino acids. Any of the amino acids
25 described above are suitable, especially the naturally-occurring amino acids found as protein components, although the amino acid typically is one bearing a side chain with a basic or acidic group, e.g., lysine, arginine or glutamic acid, or a neutral group such as glycine, serine, threonine, alanine, isoleucine, or leucine.

Methods of Inhibition of Immuno-Modulation

30 Another aspect of the invention relates to methods of inhibiting the activity of immuno-modulators comprising the step of treating a sample suspected of containing an immuno-modulator with a composition of the '

invention.

Compositions of the invention may act as inhibitors of immune-modulation, as intermediates for such inhibitors or have other utilities as described below. The inhibitors will bind to locations on the surface or in a cavity of an immuno-modulator having a geometry unique to immuno-modulators. Compositions binding immuno-modulators may bind with varying degrees of reversibility. Those compounds binding substantially irreversibly are ideal candidates for use in this method of the invention. Once labeled, the substantially irreversibly binding compositions are useful as probes for the detection of immune-modulation. Accordingly, the invention relates to methods of detecting immune-modulation in a sample suspected of containing an immuno-modulator comprising the steps of: treating a sample suspected of containing an immuno-modulator with a composition comprising a compound of the invention bound to a label; and observing the effect of the sample on the activity of the label. Suitable labels are well known in the diagnostics field and include stable free radicals, fluorophores, radioisotopes, enzymes, chemiluminescent groups and chromogens. The compounds herein are labeled in conventional fashion using functional groups such as hydroxyl or amino.

Within the context of the invention samples suspected of containing an immuno-modulator include natural or man-made materials such as living organisms; tissue or cell cultures; biological samples such as biological material samples (blood, serum, urine, cerebrospinal fluid, tears, sputum, saliva, tissue samples, and the like); laboratory samples; food, water, or air samples; bioproduct samples such as extracts of cells, particularly recombinant cells synthesizing a desired glycoprotein; and the like. Typically the sample will be suspected of containing an immuno-modulator. Samples can be contained in any medium including water and organic solvent/water mixtures. Samples include living organisms such as humans, and man made materials such as cell cultures.

The treating step of the invention comprises adding the composition of the invention to the sample or it comprises adding a precursor of the composition to the sample. The addition step comprises any method of administration as described above.

If desired, the activity of an immuno-modulator, after application of the

composition, can be observed by any method including direct and indirect methods of detecting immuno-modulator activity. Quantitative, qualitative, and semiquantitative methods of determining immuno-modulation activity are all contemplated. Typically one of the screening methods described above are
5 applied, however, any other method such as observation of the physiological properties of a living organism are also applicable.

However, in screening compounds capable of inhibiting immuno-modulation it should be kept in mind that the results of enzyme assays may not correlate with cell culture assays. Thus, a cell based assay should be the primary
10 screening tool.

Screens for Immuno-Modulating Inhibitors

Compositions of the invention are screened for inhibitory activity against immuno-modulators by any of the conventional techniques for evaluating enzyme activity. Within the context of the invention, typically compositions are
15 first screened for inhibition of immuno-modulators *in vitro* and compositions showing inhibitory activity are then screened for activity *in vivo*. Compositions having *in vitro* K_i (inhibitory constants) of less than about 5×10^{-6} M, typically less than about 1×10^{-7} M and preferably less than about 5×10^{-8} M are preferred for *in vivo* use.

20 Useful *in vitro* screens have been described in detail and will not be elaborated here. However, the examples describe suitable *in vitro* assays.

Pharmaceutical Formulations

The compounds of this invention are formulated with conventional carriers and excipients, which will be selected in accord with ordinary practice.
25 Tablets will contain excipients, glidants, fillers, binders and the like. Aqueous formulations are prepared in sterile form, and when intended for delivery by other than oral administration generally will be isotonic. All formulations will optionally contain excipients such as those set forth in the Handbook of Pharmaceutical Excipients (1986). Excipients include ascorbic acid and other
30 antioxidants, chelating agents such as EDTA, carbohydrates such as dextrin, hydroxyalkylcellulose, hydroxyalkylmethylcellulose, stearic acid and the like. The pH of the formulations ranges from about 3 to about 11, but is ordinarily

about 7 to 10.

While it is possible for the active ingredients to be administered alone it may be preferable to present them as pharmaceutical formulations. The formulations, both for veterinary and for human use, of the invention comprise at least one active ingredient, as above defined, together with one or more acceptable carriers therefor and optionally other therapeutic ingredients. The carrier(s) must be "acceptable" in the sense of being compatible with the other ingredients of the formulation and physiologically innocuous to the recipient thereof.

10 The formulations include those suitable for the foregoing administration routes. The formulations may conveniently be presented in unit dosage form and may be prepared by any of the methods well known in the art of pharmacy. Techniques and formulations generally are found in Remington's Pharmaceutical Sciences (Mack Publishing Co., Easton, PA). Such methods include the step of bringing into association the active ingredient with the carrier which constitutes one or more accessory ingredients. In general the formulations are prepared by uniformly and intimately bringing into association the active ingredient with liquid carriers or finely divided solid carriers or both, and then, if necessary, shaping the product.

20 Formulations of the present invention suitable for oral administration may be presented as discrete units such as capsules, cachets or tablets each containing a predetermined amount of the active ingredient; as a powder or granules; as a solution or a suspension in an aqueous or non-aqueous liquid; or as an oil-in-water liquid emulsion or a water-in-oil liquid emulsion. The active ingredient may also be administered as a bolus, electuary or paste.

 A tablet is made by compression or molding, optionally with one or more accessory ingredients. Compressed tablets may be prepared by compressing in a suitable machine the active ingredient in a free-flowing form such as a powder or granules, optionally mixed with a binder, lubricant, inert diluent, preservative, surface active or dispersing agent. Molded tablets may be made by molding in a suitable machine a mixture of the powdered active ingredient moistened with an inert liquid diluent. The tablets may optionally be coated or scored and optionally are formulated so as to provide slow or controlled release of the active

ingredient therefrom.

For administration to the eye or other external tissues e.g., mouth and skin, the formulations are preferably applied as a topical ointment or cream containing the active ingredient(s) in an amount of, for example, 0.075 to 20% w/w (including active ingredient(s) in a range between 0.1% and 20% in increments of 0.1% w/w such as 0.6% w/w, 0.7% w/w, etc.), preferably 0.2 to 15% w/w and most preferably 0.5 to 10% w/w. When formulated in an ointment, the active ingredients may be employed with either a paraffinic or a water-miscible ointment base. Alternatively, the active ingredients may be formulated in a cream with an oil-in-water cream base.

If desired, the aqueous phase of the cream base may include, for example, at least 30% w/w of a polyhydric alcohol, i.e. an alcohol having two or more hydroxyl groups such as propylene glycol, butane 1,3-diol, mannitol, sorbitol, glycerol and polyethylene glycol (including PEG 400) and mixtures thereof. The topical formulations may desirably include a compound which enhances absorption or penetration of the active ingredient through the skin or other affected areas. Examples of such dermal penetration enhancers include dimethyl sulphoxide and related analogs.

The oily phase of the emulsions of this invention may be constituted from known ingredients in a known manner. While the phase may comprise merely an emulsifier (otherwise known as an emulgent), it desirably comprises a mixture of at least one emulsifier with a fat or an oil or with both a fat and an oil. Preferably, a hydrophilic emulsifier is included together with a lipophilic emulsifier which acts as a stabilizer. It is also preferred to include both an oil and a fat. Together, the emulsifier(s) with or without stabilizer(s) make up the so-called emulsifying wax, and the wax together with the oil and fat make up the so-called emulsifying ointment base which forms the oily dispersed phase of the cream formulations.

Emulgents and emulsion stabilizers suitable for use in the formulation of the invention include Tween[®] 60, Span[®] 80, cetostearyl alcohol, benzyl alcohol, myristyl alcohol, glyceryl mono-stearate and sodium lauryl sulfate.

The choice of suitable oils or fats for the formulation is based on achieving the desired cosmetic properties. The cream should preferably be a

non-greasy, non-staining and washable product with suitable consistency to avoid leakage from tubes or other containers. Straight or branched chain, mono- or dibasic alkyl esters such as di-isoadipate, isocetyl stearate, propylene glycol diester of coconut fatty acids, isopropyl myristate, decyl oleate, isopropyl palmitate, butyl stearate, 2-ethylhexyl palmitate or a blend of branched chain esters known as Crodamol CAP may be used, the last three being preferred esters. These may be used alone or in combination depending on the properties required. Alternatively, high melting point lipids such as white soft paraffin and/or liquid paraffin or other mineral oils are used.

Pharmaceutical formulations according to the present invention comprise one or more compounds of the invention together with one or more pharmaceutically acceptable carriers or excipients and optionally other therapeutic agents. Pharmaceutical formulations containing the active ingredient may be in any form suitable for the intended method of administration. When used for oral use for example, tablets, troches, lozenges, aqueous or oil suspensions, dispersible powders or granules, emulsions, hard or soft capsules, syrups or elixirs may be prepared. Compositions intended for oral use may be prepared according to any method known to the art for the manufacture of pharmaceutical compositions and such compositions may contain one or more agents including sweetening agents, flavoring agents, coloring agents and preserving agents, in order to provide a palatable preparation. Tablets containing the active ingredient in admixture with non-toxic pharmaceutically acceptable excipient which are suitable for manufacture of tablets are acceptable. These excipients may be, for example, inert diluents, such as calcium or sodium carbonate, lactose, lactose monohydrate, croscarmellose sodium, povidone, calcium or sodium phosphate; granulating and disintegrating agents, such as maize starch, or alginic acid; binding agents, such as cellulose, microcrystalline cellulose, starch, gelatin or acacia; and lubricating agents, such as magnesium stearate, stearic acid or talc. Tablets may be uncoated or may be coated by known techniques including microencapsulation to delay disintegration and adsorption in the gastrointestinal tract and thereby provide a sustained action over a longer period. For example, a time delay material such as glyceryl monostearate or glyceryl distearate alone or with a wax may be employed.

Formulations for oral use may be also presented as hard gelatin capsules where the active ingredient is mixed with an inert solid diluent, for example calcium phosphate or kaolin, or as soft gelatin capsules wherein the active ingredient is mixed with water or an oil medium, such as peanut oil, liquid paraffin or olive oil.

Aqueous suspensions of the invention contain the active materials in admixture with excipients suitable for the manufacture of aqueous suspensions. Such excipients include a suspending agent, such as sodium carboxymethylcellulose, methylcellulose, hydroxypropyl methylcellulose, sodium alginate, polyvinylpyrrolidone, gum tragacanth and gum acacia, and dispersing or wetting agents such as a naturally occurring phosphatide (e.g., lecithin), a condensation product of an alkylene oxide with a fatty acid (e.g., polyoxyethylene stearate), a condensation product of ethylene oxide with a long chain aliphatic alcohol (e.g., heptadecaethyleneoxycetanol), a condensation product of ethylene oxide with a partial ester derived from a fatty acid and a hexitol anhydride (e.g., polyoxyethylene sorbitan monooleate). The aqueous suspension may also contain one or more preservatives such as ethyl or n-propyl p-hydroxy-benzoate, one or more coloring agents, one or more flavoring agents and one or more sweetening agents, such as sucrose or saccharin.

Oil suspensions may be formulated by suspending the active ingredient in a vegetable oil, such as arachis oil, olive oil, sesame oil or coconut oil, or in a mineral oil such as liquid paraffin. The oral suspensions may contain a thickening agent, such as beeswax, hard paraffin or cetyl alcohol. Sweetening agents, such as those set forth above, and flavoring agents may be added to provide a palatable oral preparation. These compositions may be preserved by the addition of an antioxidant such as ascorbic acid.

Dispersible powders and granules of the invention suitable for preparation of an aqueous suspension by the addition of water provide the active ingredient in admixture with a dispersing or wetting agent, a suspending agent, and one or more preservatives. Suitable dispersing or wetting agents and suspending agents are exemplified by those disclosed above. Additional excipients, for example sweetening, flavoring and coloring agents, may also be present.

The pharmaceutical compositions of the invention may also be in the form of oil-in-water emulsions. The oily phase may be a vegetable oil, such as olive oil or arachis oil, a mineral oil, such as liquid paraffin, or a mixture of these. Suitable emulsifying agents include naturally-occurring gums, such as gum acacia and gum tragacanth, naturally occurring phosphatides, such as soybean lecithin, esters or partial esters derived from fatty acids and hexitol anhydrides, such as sorbitan monooleate, and condensation products of these partial esters with ethylene oxide, such as polyoxyethylene sorbitan monooleate. The emulsion may also contain sweetening and flavoring agents. Syrups and elixirs may be formulated with sweetening agents, such as glycerol, sorbitol or sucrose. Such formulations may also contain a demulcent, a preservative, a flavoring or a coloring agent.

The pharmaceutical compositions of the invention may be in the form of a sterile injectable preparation, such as a sterile injectable aqueous or oleaginous suspension. This suspension may be formulated according to the known art using those suitable dispersing or wetting agents and suspending agents which have been mentioned above. The sterile injectable preparation may also be a sterile injectable solution or suspension in a non-toxic parenterally acceptable diluent or solvent, such as a solution in 1,3-butane-diol or prepared as a lyophilized powder. Among the acceptable vehicles and solvents that may be employed are water, Ringer's solution and isotonic sodium chloride solution. In addition, sterile fixed oils may conventionally be employed as a solvent or suspending medium. For this purpose any bland fixed oil may be employed including synthetic mono- or diglycerides. In addition, fatty acids such as oleic acid may likewise be used in the preparation of injectables.

The amount of active ingredient that may be combined with the carrier material to produce a single dosage form will vary depending upon the host treated and the particular mode of administration. For example, a time-release formulation intended for oral administration to humans may contain approximately 1 to 1000 mg of active material compounded with an appropriate and convenient amount of carrier material which may vary from about 5 to about 95% of the total compositions (weight:weight). The pharmaceutical composition can be prepared to provide easily measurable amounts for administration. For

example, an aqueous solution intended for intravenous infusion may contain from about 3 to 500 μg of the active ingredient per milliliter of solution in order that infusion of a suitable volume at a rate of about 30 mL/hr can occur.

Formulations suitable for administration to the eye include eye drops
5 wherein the active ingredient is dissolved or suspended in a suitable carrier, especially an aqueous solvent for the active ingredient. The active ingredient is preferably present in such formulations in a concentration of 0.5 to 20%, advantageously 0.5 to 10% particularly about 1.5% w/w.

Formulations suitable for topical administration in the mouth include
10 lozenges comprising the active ingredient in a flavored basis, usually sucrose and acacia or tragacanth; pastilles comprising the active ingredient in an inert basis such as gelatin and glycerin, or sucrose and acacia; and mouthwashes comprising the active ingredient in a suitable liquid carrier.

Formulations for rectal administration may be presented as a suppository
15 with a suitable base comprising for example cocoa butter or a salicylate.

Formulations suitable for intrapulmonary or nasal administration have a particle size for example in the range of 0.1 to 500 microns (including particle sizes in a range between 0.1 and 500 microns in increments microns such as 0.5, 1, 30 microns, 35 microns, etc.), which is administered by rapid inhalation
20 through the nasal passage or by inhalation through the mouth so as to reach the alveolar sacs. Suitable formulations include aqueous or oily solutions of the active ingredient. Formulations suitable for aerosol or dry powder administration may be prepared according to conventional methods and may be delivered with other therapeutic agents such as compounds heretofore used in the
25 treatment or prophylaxis of infections as described below.

Formulations suitable for vaginal administration may be presented as pessaries, tampons, creams, gels, pastes, foams or spray formulations containing in addition to the active ingredient such carriers as are known in the art to be appropriate.

30 Formulations suitable for parenteral administration include aqueous and non-aqueous sterile injection solutions which may contain anti-oxidants, buffers, bacteriostats and solutes which render the formulation isotonic with the blood of the intended recipient; and aqueous and non-aqueous sterile suspensions which

may include suspending agents and thickening agents.

The formulations are presented in unit-dose or multi-dose containers, for example sealed ampoules and vials, and may be stored in a freeze-dried (lyophilized) condition requiring only the addition of the sterile liquid carrier, for example water for injection, immediately prior to use. Extemporaneous injection solutions and suspensions are prepared from sterile powders, granules and tablets of the kind previously described. Preferred unit dosage formulations are those containing a daily dose or unit daily sub-dose, as herein above recited, or an appropriate fraction thereof, of the active ingredient.

It should be understood that in addition to the ingredients particularly mentioned above the formulations of this invention may include other agents conventional in the art having regard to the type of formulation in question, for example those suitable for oral administration may include flavoring agents.

The invention further provides veterinary compositions comprising at least one active ingredient as above defined together with a veterinary carrier therefor.

Veterinary carriers are materials useful for the purpose of administering the composition and may be solid, liquid or gaseous materials which are otherwise inert or acceptable in the veterinary art and are compatible with the active ingredient. These veterinary compositions may be administered orally, parenterally or by any other desired route.

Compounds of the invention can also be formulated to provide controlled release of the active ingredient to allow less frequent dosing or to improve the pharmacokinetic or toxicity profile of the active ingredient. Accordingly, the invention also provided compositions comprising one or more compounds of the invention formulated for sustained or controlled release.

Effective dose of active ingredient depends at least on the nature of the condition being treated, toxicity, whether the compound is being used prophylactically (lower doses) or against an active infection, the method of delivery, and the pharmaceutical formulation, and will be determined by the clinician using conventional dose escalation studies. It can be expected to be from about 0.0001 to about 100 mg/kg body weight per day. Typically, from about 0.01 to about 10 mg/kg body weight per day. More typically, from about

.01 to about 5 mg/kg body weight per day. More typically, from about .05 to about 0.5 mg/kg body weight per day. For example, the daily candidate dose for an adult human of approximately 70 kg body weight will range from 1 mg to 1000 mg, preferably between 5 mg and 500 mg, and may take the form of single or multiple doses.

Routes of Administration

One or more compounds of the invention (herein referred to as the active ingredients) are administered by any route appropriate to the condition to be treated. Suitable routes include oral, rectal, nasal, topical (including buccal and sublingual), vaginal and parenteral (including subcutaneous, intramuscular, intravenous, intradermal, intrathecal and epidural), and the like. It will be appreciated that the preferred route may vary with for example the condition of the recipient. An advantage of the compounds of this invention is that they are orally bioavailable and can be dosed orally.

Combination Therapy

Active ingredients of the invention are also used in combination with other active ingredients. Such combinations are selected based on the condition to be treated, cross-reactivities of ingredients and pharmaco-properties of the combination.

It is also possible to combine any compound of the invention with one or more other active ingredients in a unitary dosage form for simultaneous or sequential administration to a patient. The combination therapy may be administered as a simultaneous or sequential regimen. When administered sequentially, the combination may be administered in two or more administrations.

The combination therapy may provide "synergy" and "synergistic effect", i.e. the effect achieved when the active ingredients used together is greater than the sum of the effects that results from using the compounds separately. A synergistic effect may be attained when the active ingredients are: (1) co-formulated and administered or delivered simultaneously in a combined formulation; (2) delivered by alternation or in parallel as separate formulations; or (3) by some other regimen. When delivered in alternation therapy, a

synergistic effect may be attained when the compounds are administered or delivered sequentially, e.g., in separate tablets, pills or capsules, or by different injections in separate syringes. In general, during alternation therapy, an effective dosage of each active ingredient is administered sequentially, i.e.
5 serially, whereas in combination therapy, effective dosages of two or more active ingredients are administered together.

Metabolites of the Compounds of the Invention

Also falling within the scope of this invention are the *in vivo* metabolic products of the compounds described herein. Such products may result for
10 example from the oxidation, reduction, hydrolysis, amidation, esterification and the like of the administered compound, primarily due to enzymatic processes. Accordingly, the invention includes compounds produced by a process comprising contacting a compound of this invention with a mammal for a period of time sufficient to yield a metabolic product thereof. Such products typically
15 are identified by preparing a radiolabelled (e.g., C¹⁴ or H³) compound of the invention, administering it parenterally in a detectable dose (e.g., greater than about 0.5 mg/kg) to an animal such as rat, mouse, guinea pig, monkey, or to man, allowing sufficient time for metabolism to occur (typically about 30 seconds to 30 hours) and isolating its conversion products from the urine, blood
20 or other biological samples. These products are easily isolated since they are labeled (others are isolated by the use of antibodies capable of binding epitopes surviving in the metabolite). The metabolite structures are determined in conventional fashion, e.g., by MS or NMR analysis. In general, analysis of metabolites is done in the same way as conventional drug metabolism studies
25 well-known to those skilled in the art. The conversion products, so long as they are not otherwise found *in vivo*, are useful in diagnostic assays for therapeutic dosing of the compounds of the invention even if they possess no immuno-modulation inhibitory activity of their own.

Recipes and methods for determining stability of compounds in surrogate
30 gastrointestinal secretions are known. Compounds are defined herein as stable in the gastrointestinal tract where less than about 50 mole percent of the protected groups are deprotected in surrogate intestinal or gastric juice upon

incubation for 1 hour at 37 °C. Simply because the compounds are stable to the gastrointestinal tract does not mean that they cannot be hydrolyzed *in vivo*. The phosphonate prodrugs of the invention typically will be stable in the digestive system but are substantially hydrolyzed to the parental drug in the digestive
5 lumen, liver or other metabolic organ, or within cells in general.

Exemplary Methods of Making the Compounds of the Invention.

The invention also relates to methods of making the compositions of the invention. The compositions are prepared by any of the applicable techniques of organic synthesis. Many such techniques are well known in the art. However,
10 many of the known techniques are elaborated in Compendium of Organic Synthetic Methods (John Wiley & Sons, New York), Vol. 1, Ian T. Harrison and Shuyen Harrison, 1971; Vol. 2, Ian T. Harrison and Shuyen Harrison, 1974; Vol. 3, Louis S. Hegedus and Leroy Wade, 1977; Vol. 4, Leroy G. Wade, jr., 1980; Vol. 5, Leroy G. Wade, Jr., 1984; and Vol. 6, Michael B. Smith; as well as
15 March, J., Advanced Organic Chemistry, Third Edition, (John Wiley & Sons, New York, 1985), Comprehensive Organic Synthesis. Selectivity, Strategy & Efficiency in Modern Organic Chemistry. In 9 Volumes, Barry M. Trost, Editor-in-Chief (Pergamon Press, New York, 1993 printing).

A number of exemplary methods for the preparation of the compositions
20 of the invention are provided below. These methods are intended to illustrate the nature of such preparations are not intended to limit the scope of applicable methods.

Generally, the reaction conditions such as temperature, reaction time, solvents, work-up procedures, and the like, will be those common in the art for
25 the particular reaction to be performed. The cited reference material, together with material cited therein, contains detailed descriptions of such conditions. Typically the temperatures will be -100°C to 200°C, solvents will be aprotic or protic, and reaction times will be 10 seconds to 10 days. Work-up typically consists of quenching any unreacted reagents followed by partition between a
30 water/organic layer system (extraction) and separating the layer containing the product.

Oxidation and reduction reactions are typically carried out at temperatures near room temperature (about 20 °C), although for metal hydride

reductions frequently the temperature is reduced to 0 °C to -100 °C, solvents are typically aprotic for reductions and may be either protic or aprotic for oxidations. Reaction times are adjusted to achieve desired conversions.

Condensation reactions are typically carried out at temperatures near
5 room temperature, although for non-equilibrating, kinetically controlled condensations reduced temperatures (0 °C to -100 °C) are also common. Solvents can be either protic (common in equilibrating reactions) or aprotic (common in kinetically controlled reactions).

Standard synthetic techniques such as azeotropic removal of reaction by-
10 products and use of anhydrous reaction conditions (e.g., inert gas environments) are common in the art and will be applied when applicable.

Schemes and Examples

General aspects of these exemplary methods are described below and in the Examples. Each of the products of the following processes is optionally
15 separated, isolated, and/or purified prior to its use in subsequent processes.

Generally, the reaction conditions such as temperature, reaction time, solvents, work-up procedures, and the like, will be those common in the art for the particular reaction to be performed. The cited reference material, together with material cited therein, contains detailed descriptions of such conditions.
20 Typically the temperatures will be -100°C to 200°C, solvents will be aprotic or protic, and reaction times will be 10 seconds to 10 days. Work-up typically consists of quenching any unreacted reagents followed by partition between a water/organic layer system (extraction) and separating the layer containing the product.

25 Oxidation and reduction reactions are typically carried out at temperatures near room temperature (about 20 °C), although for metal hydride reductions frequently the temperature is reduced to 0 °C to -100 °C, solvents are typically aprotic for reductions and may be either protic or aprotic for oxidations. Reaction times are adjusted to achieve desired conversions.

30 Condensation reactions are typically carried out at temperatures near room temperature, although for non-equilibrating, kinetically controlled condensations reduced temperatures (0 °C to -100 °C) are also common. Solvents can be either protic (common in equilibrating reactions) or aprotic

(common in kinetically controlled reactions).

Standard synthetic techniques such as azeotropic removal of reaction by-products and use of anhydrous reaction conditions (e.g., inert gas environments) are common in the art and will be applied when applicable.

5 The terms “treated”, “treating”, “treatment”, and the like, when used in connection with a chemical synthetic operation, mean contacting, mixing, reacting, allowing to react, bringing into contact, and other terms common in the art for indicating that one or more chemical entities is treated in such a manner as to convert it to one or more other chemical entities. This means that “treating
10 compound one with compound two” is synonymous with “allowing compound one to react with compound two”, “contacting compound one with compound two”, “reacting compound one with compound two”, and other expressions common in the art of organic synthesis for reasonably indicating that compound one was “treated”, “reacted”, “allowed to react”, etc., with compound two. For
15 example, treating indicates the reasonable and usual manner in which organic chemicals are allowed to react. Normal concentrations (0.01M to 10M, typically 0.1M to 1M), temperatures (-100 °C to 250 °C, typically -78 °C to 150 °C, more typically -78 °C to 100 °C, still more typically 0 °C to 100 °C), reaction vessels (typically glass, plastic, metal), solvents, pressures, atmospheres (typically air
20 for oxygen and water insensitive reactions or nitrogen or argon for oxygen or water sensitive), etc., are intended unless otherwise indicated. The knowledge of similar reactions known in the art of organic synthesis are used in selecting the conditions and apparatus for “treating” in a given process. In particular, one of ordinary skill in the art of organic synthesis selects conditions and apparatus
25 reasonably expected to successfully carry out the chemical reactions of the described processes based on the knowledge in the art.

 Modifications of each of the exemplary schemes and in the examples (hereafter “exemplary schemes”) leads to various analogs of the specific exemplary materials produce. The above-cited citations describing suitable
30 methods of organic synthesis are applicable to such modifications.

 In each of the exemplary schemes it may be advantageous to separate reaction products from one another and/or from starting materials. The desired products of each step or series of steps is separated and/or purified (hereinafter

separated) to the desired degree of homogeneity by the techniques common in the art. Typically such separations involve multiphase extraction, crystallization from a solvent or solvent mixture, distillation, sublimation, or chromatography. Chromatography can involve any number of methods including, for example:

5 reverse-phase and normal phase; size exclusion; ion exchange; high, medium, and low pressure liquid chromatography methods and apparatus; small scale analytical; simulated moving bed (SMB) and preparative thin or thick layer chromatography, as well as techniques of small scale thin layer and flash chromatography.

10 Another class of separation methods involves treatment of a mixture with a reagent selected to bind to or render otherwise separable a desired product, unreacted starting material, reaction by product, or the like. Such reagents include adsorbents or absorbents such as activated carbon, molecular sieves, ion exchange media, or the like. Alternatively, the reagents can be acids in the case

15 of a basic material, bases in the case of an acidic material, binding reagents such as antibodies, binding proteins, selective chelators such as crown ethers, liquid/liquid ion extraction reagents (LIX), or the like.

Selection of appropriate methods of separation depends on the nature of the materials involved. For example, boiling point, and molecular weight in

20 distillation and sublimation, presence or absence of polar functional groups in chromatography, stability of materials in acidic and basic media in multiphase extraction, and the like. One skilled in the art will apply techniques most likely to achieve the desired separation.

A single stereoisomer, e.g., an enantiomer, substantially free of its

25 stereoisomer may be obtained by resolution of the racemic mixture using a method such as formation of diastereomers using optically active resolving agents (Stereochemistry of Carbon Compounds, (1962) by E. L. Eliel, McGraw Hill; Lochmuller, C. H., (1975) *J. Chromatogr.*, 113:(3) 283-302). Racemic mixtures of chiral compounds of the invention can be separated and isolated by

30 any suitable method, including: (1) formation of ionic, diastereomeric salts with chiral compounds and separation by fractional crystallization or other methods, (2) formation of diastereomeric compounds with chiral derivatizing reagents, separation of the diastereomers, and conversion to the pure stereoisomers, and

(3) separation of the substantially pure or enriched stereoisomers directly under chiral conditions.

Under method (1), diastereomeric salts can be formed by reaction of enantiomerically pure chiral bases such as brucine, quinine, ephedrine, 5 strychnine, α -methyl- β -phenylethylamine (amphetamine), and the like with asymmetric compounds bearing acidic functionality, such as carboxylic acid and sulfonic acid. The diastereomeric salts may be induced to separate by fractional crystallization or ionic chromatography. For separation of the optical isomers of amino compounds, addition of chiral carboxylic or sulfonic acids, such as 10 camphorsulfonic acid, tartaric acid, mandelic acid, or lactic acid can result in formation of the diastereomeric salts.

Alternatively, by method (2), the substrate to be resolved is reacted with one enantiomer of a chiral compound to form a diastereomeric pair (Eliel, E. and Wilen, S. (1994) Stereochemistry of Organic Compounds, John Wiley & Sons, 15 Inc., p. 322). Diastereomeric compounds can be formed by reacting asymmetric compounds with enantiomerically pure chiral derivatizing reagents, such as menthyl derivatives, followed by separation of the diastereomers and hydrolysis to yield the free, enantiomerically enriched xanthene. A method of determining optical purity involves making chiral esters, such as a menthyl ester, e.g., (-) 20 menthyl chloroformate in the presence of base, or Mosher ester, α -methoxy- α -(trifluoromethyl)phenyl acetate (Jacob III. (1982) *J. Org. Chem.* 47:4165), of the racemic mixture, and analyzing the NMR spectrum for the presence of the two atropisomeric diastereomers. Stable diastereomers of atropisomeric compounds can be separated and isolated by normal- and reverse-phase chromatography 25 following methods for separation of atropisomeric naphthyl-isoquinolines (Hoye, T., WO 96/15111). By method (3), a racemic mixture of two enantiomers can be separated by chromatography using a chiral stationary phase (Chiral Liquid Chromatography (1989) W. J. Lough, Ed. Chapman and Hall, New York; Okamoto, (1990) *J. of Chromatogr.* 513:375-378). Enriched or 30 purified enantiomers can be distinguished by methods used to distinguish other chiral molecules with asymmetric carbon atoms, such as optical rotation and circular dichroism.

Examples General Section

A number of exemplary methods for the preparation of compounds of the invention are provided herein, for example, in the Examples hereinbelow..

These methods are intended to illustrate the nature of such preparations are not
5 intended to limit the scope of applicable methods. Certain compounds of the invention can be used as intermediates for the preparation of other compounds of the invention. For example, the interconversion of various phosphonate compounds of the invention is illustrated below.

10 INTERCONVERSIONS OF THE PHOSPHONATES R-LINK-P(O)(OR¹)₂, R-LINK-P(O)(OR¹)(OH) AND R-LINK-P(O)(OH)₂.

The following schemes 32-38 described the preparation of phosphonate esters of the general structure R-link-P(O)(OR¹)₂, in which the groups R¹ may be the same or different. The R¹ groups attached to a phosphonate ester, or to
15 precursors thereto, may be changed using established chemical transformations. The interconversion reactions of phosphonates are illustrated in Scheme S32. The group R in Scheme 32 represents the substructure, i.e. the drug "scaffold, to which the substituent link-P(O)(OR¹)₂ is attached, either in the compounds of the invention, or in precursors thereto. At the point in the synthetic route of
20 conducting a phosphonate interconversion, certain functional groups in R may be protected. The methods employed for a given phosphonate transformation depend on the nature of the substituent R¹, and of the substrate to which the phosphonate group is attached. The preparation and hydrolysis of phosphonate esters is described in Organic Phosphorus Compounds, G. M. Kosolapoff, L.
25 Maeir, eds, Wiley, 1976, p. 9ff.

In general, synthesis of phosphonate esters is achieved by coupling a nucleophile amine or alcohol with the corresponding activated phosphonate electrophilic precursor. For example, chlorophosphonate addition on to 5'-hydroxy of nucleoside is a well known method for preparation of nucleoside
30 phosphate monoesters. The activated precursor can be prepared by several well known methods. Chlorophosphonates useful for synthesis of the prodrugs are prepared from the substituted-1,3-propanediol (Wissner, et al, (1992) *J. Med Chem.* 35:1650). Chlorophosphonates are made by oxidation of the

corresponding chlorophospholanes (Anderson, et al, (1984) *J. Org. Chem.* 49:1304) which are obtained by reaction of the substituted diol with phosphorus trichloride. Alternatively, the chlorophosphonate agent is made by treating substituted-1,3-diols with phosphorusoxychloride (Patois, et al, (1990) *J. Chem. Soc. Perkin Trans. I*, 1577). Chlorophosphonate species may also be generated in situ from corresponding cyclic phosphites (Silverburg, et al., (1996) *Tetrahedron lett.*, 37:771-774), which in turn can be either made from chlorophospholane or phosphoramidate intermediate. Phosphorofluoridate intermediate prepared either from pyrophosphate or phosphoric acid may also act as precursor in preparation of cyclic prodrugs (Watanabe et al., (1988) *Tetrahedron lett.*, 29:5763-66).

Phosphonate prodrugs of the present invention may also be prepared from the free acid by Mitsunobu reactions (Mitsunobu, (1981) *Synthesis*, 1; Campbell, (1992) *J. Org. Chem.* 57:6331), and other acid coupling reagents including, but not limited to, carbodiimides (Alexander, et al, (1994) *Collect. Czech. Chem. Commun.* 59:1853; Casara et al, (1992) *Bioorg. Med. Chem. Lett.* 2:145; Ohashi et al, (1988) *Tetrahedron Lett.*, 29:1189), and benzotriazolyloxytris-(dimethylamino)phosphonium salts (Campagne et al (1993) *Tetrahedron Lett.* 34:6743).

Aryl halides undergo Ni^{+2} catalyzed reaction with phosphite derivatives to give aryl phosphonate containing compounds (Balthazar, et al (1980) *J. Org. Chem.* 45:5425). Phosphonates may also be prepared from the chlorophosphonate in the presence of a palladium catalyst using aromatic triflates (Petrakis et al (1987) *J. Am. Chem. Soc.* 109:2831; Lu et al (1987) *Synthesis* 726). In another method, aryl phosphonate esters are prepared from aryl phosphates under anionic rearrangement conditions (Melvin (1981) *Tetrahedron Lett.* 22:3375; Casteel et al (1991) *Synthesis*, 691). N-Alkoxy aryl salts with alkali metal derivatives of cyclic alkyl phosphonate provide general synthesis for heteroaryl-2-phosphonate linkers (Redmore (1970) *J. Org. Chem.* 35:4114). These above mentioned methods can also be extended to compounds where the W^5 group is a heterocycle. Cyclic-1,3-propanyl prodrugs of phosphonates are also synthesized from phosphonic diacids and substituted propane-1,3-diols using a coupling reagent such as 1,3-dicyclohexylcarbodiimide (DCC) in presence of a base (e.g., pyridine). Other

carbodiimide based coupling agents like 1,3-disopropylcarbodiimide or water soluble reagent, 1-(3-dimethylaminopropyl)-3-ethylcarbodiimide hydrochloride (EDCI) can also be utilized for the synthesis of cyclic phosphonate prodrugs.

The conversion of a phosphonate diester S32.1 into the corresponding
5 phosphonate monoester S32.2 (Scheme 32, Reaction 1) is accomplished by a number of methods. For example, the ester S32.1 in which R¹ is an aralkyl group such as benzyl, is converted into the monoester compound S32.2 by reaction with a tertiary organic base such as diazabicyclooctane (DABCO) or quinuclidine, as described in *J. Org. Chem.* (1995) 60:2946. The reaction is
10 performed in an inert hydrocarbon solvent such as toluene or xylene, at about 110 °C. The conversion of the diester S32.1 in which R¹ is an aryl group such as phenyl, or an alkenyl group such as allyl, into the monoester S32.2 is effected by treatment of the ester S32.1 with a base such as aqueous sodium hydroxide in acetonitrile or lithium hydroxide in aqueous tetrahydrofuran. Phosphonate
15 diesters S32.1 in which one of the groups R¹ is aralkyl, such as benzyl, and the other is alkyl, is converted into the monoesters S32.2 in which R¹ is alkyl by hydrogenation, for example using a palladium on carbon catalyst. Phosphonate diesters in which both of the groups R¹ are alkenyl, such as allyl, is converted into the monoester S32.2 in which R¹ is alkenyl, by treatment with
20 chlorotris(triphenylphosphine)rhodium (Wilkinson's catalyst) in aqueous ethanol at reflux, optionally in the presence of diazabicyclooctane, for example by using the procedure described in *J. Org. Chem.* (1973) 38:3224, for the cleavage of allyl carboxylates.

The conversion of a phosphonate diester S32.1 or a phosphonate
25 monoester S32.2 into the corresponding phosphonic acid S32.3 (Scheme 32, Reactions 2 and 3) can be effected by reaction of the diester or the monoester with trimethylsilyl bromide, as described in *J. Chem. Soc., Chem. Comm.*, (1979) 739. The reaction is conducted in an inert solvent such as, for example, dichloromethane, optionally in the presence of a silylating agent such as
30 bis(trimethylsilyl)trifluoroacetamide, at ambient temperature. A phosphonate monoester S32.2 in which R¹ is aralkyl such as benzyl, is converted into the corresponding phosphonic acid S32.3 by hydrogenation over a palladium catalyst, or by treatment with hydrogen chloride in an ethereal solvent such as

dioxane. A phosphonate monoester S32.2 in which R¹ is alkenyl such as, for example, allyl, is converted into the phosphonic acid S32.3 by reaction with Wilkinson's catalyst in an aqueous organic solvent, for example in 15% aqueous acetonitrile, or in aqueous ethanol, for example using the procedure described in
5 *Helv. Chim. Acta.* (1985) 68:618. Palladium catalyzed hydrogenolysis of phosphonate esters S32.1 in which R¹ is benzyl is described in *J. Org. Chem.* (1959) 24:434. Platinum-catalyzed hydrogenolysis of phosphonate esters S32.1 in which R¹ is phenyl is described in *J. Am. Chem. Soc.* (1956) 78:2336.

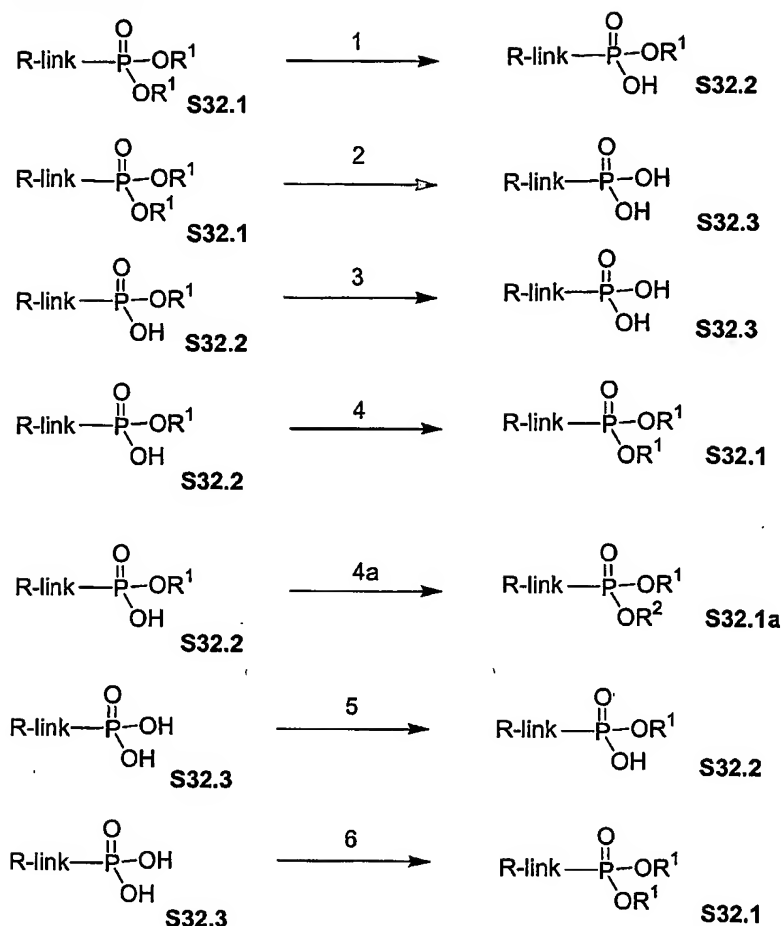
The conversion of a phosphonate monoester S32.2 into a phosphonate
10 diester S32.1 (Scheme 32, Reaction 4) in which the newly introduced R¹ group is alkyl, aralkyl, haloalkyl such as chloroethyl, or aralkyl is effected by a number of reactions in which the substrate S32.2 is reacted with a hydroxy compound R¹OH, in the presence of a coupling agent. Typically, the second phosphonate ester group is different than the first introduced phosphonate ester group, i.e. R¹
15 is followed by the introduction of R² where each of R¹ and R² is alkyl, aralkyl, haloalkyl such as chloroethyl, or aralkyl (Scheme 32, Reaction 4a) whereby S32.2 is converted to S32.1a. Suitable coupling agents are those employed for the preparation of carboxylate esters, and include a carbodiimide such as dicyclohexylcarbodiimide, in which case the reaction is preferably conducted in
20 a basic organic solvent such as pyridine, or (benzotriazol-1-yl)oxytripyrrolidinophosphonium hexafluorophosphate (PYBOP, Sigma), in which case the reaction is performed in a polar solvent such as dimethylformamide, in the presence of a tertiary organic base such as diisopropylethylamine, or Aldrithiol-2 (Aldrich) in which case the reaction is
25 conducted in a basic solvent such as pyridine, in the presence of a triaryl phosphine such as triphenylphosphine. Alternatively, the conversion of the phosphonate monoester S32.2 to the diester S32.1 is effected by the use of the Mitsunobu reaction, as described above. The substrate is reacted with the hydroxy compound R¹OH, in the presence of diethyl azodicarboxylate and a
30 triarylphosphine such as triphenyl phosphine. Alternatively, the phosphonate monoester S32.2 is transformed into the phosphonate diester S32.1, in which the introduced R¹ group is alkenyl or aralkyl, by reaction of the monoester with the halide R¹Br, in which R¹ is as alkenyl or aralkyl. The alkylation reaction is

conducted in a polar organic solvent such as dimethylformamide or acetonitrile, in the presence of a base such as cesium carbonate. Alternatively, the phosphonate monoester is transformed into the phosphonate diester in a two step procedure. In the first step, the phosphonate monoester S32.2 is transformed into the chloro analog $\text{RP}(\text{O})(\text{OR}^1)\text{Cl}$ by reaction with thionyl chloride or oxalyl chloride and the like, as described in Organic Phosphorus Compounds, G. M. Kosolapoff, L. Maeir, eds, Wiley, 1976, p. 17, and the thus-obtained product $\text{RP}(\text{O})(\text{OR}^1)\text{Cl}$ is then reacted with the hydroxy compound R^1OH , in the presence of a base such as triethylamine, to afford the phosphonate diester S32.1.

10 A phosphonic acid R-link- $\text{P}(\text{O})(\text{OH})_2$ is transformed into a phosphonate monoester $\text{RP}(\text{O})(\text{OR}^1)(\text{OH})$ (Scheme 32, Reaction 5) by means of the methods described above of for the preparation of the phosphonate diester R-link- $\text{P}(\text{O})(\text{OR}^1)_2$ S32.1, except that only one molar proportion of the component R^1OH or R^1Br is employed. Dialkyl phosphonates may be prepared according to the methods of: Quast et al (1974) *Synthesis* 490; Stowell et al (1990) 15 *Tetrahedron Lett.* 3261; US 5663159.

A phosphonic acid R-link- $\text{P}(\text{O})(\text{OH})_2$ S32.3 is transformed into a phosphonate diester R-link- $\text{P}(\text{O})(\text{OR}^1)_2$ S32.1 (Scheme 32, Reaction 6) by a coupling reaction with the hydroxy compound R^1OH , in the presence of a coupling agent such as Aldrithiol-2 (Aldrich) and triphenylphosphine. The reaction is conducted in a basic solvent such as pyridine. Alternatively, phosphonic acids S32.3 are transformed into phosphonic esters S32.1 in which R^1 is aryl, by means of a coupling reaction employing, for example, dicyclohexylcarbodiimide in pyridine at ca 70 °C. Alternatively, phosphonic acids S32.3 are transformed into phosphonic esters S32.1 in which R^1 is alkenyl, by means of an alkylation reaction. The phosphonic acid is reacted with the alkenyl bromide R^1Br in a polar organic solvent such as acetonitrile solution at reflux temperature, the presence of a base such as cesium carbonate, to afford the phosphonic ester S32.1.

Scheme 32



Preparation of phosphonate carbamates.

Phosphonate esters may contain a carbamate linkage. The preparation of carbamates is described in Comprehensive Organic Functional Group Transformations, A. R. Katritzky, ed., Pergamon, 1995, Vol. 6, p. 416ff, and in Organic Functional Group Preparations, by S. R. Sandler and W. Karo, Academic Press, 1986, p. 260ff. The carbamoyl group may be formed by reaction of a hydroxy group according to the methods known in the art, including the teachings of Ellis, US 2002/0103378 A1 and Hajima, US 6018049.

Scheme 33 illustrates various methods by which the carbamate linkage is synthesized. As shown in Scheme 33, in the general reaction generating carbamates, an alcohol S33.1, is converted into the activated derivative S33.2 in which Lv is a leaving group such as halo, imidazolyl, benztriazolyl and the like, as described herein. The activated derivative S33.2 is then reacted with an amine

S33.3, to afford the carbamate product S33.4. Examples 1 – 7 in Scheme 33 depict methods by which the general reaction is effected. Examples 8 - 10 illustrate alternative methods for the preparation of carbamates.

Scheme 33, Example 1 illustrates the preparation of carbamates employing a chloroformyl derivative of the alcohol S33.5. In this procedure, the alcohol S33.5 is reacted with phosgene, in an inert solvent such as toluene, at about 0 °C, as described in *Org. Syn. Coll. Vol. 3*, 167, 1965, or with an equivalent reagent such as trichloromethoxy chloroformate, as described in *Org. Syn. Coll. Vol. 6*, 715, 1988, to afford the chloroformate S33.6. The latter compound is then reacted with the amine component S33.3, in the presence of an organic or inorganic base, to afford the carbamate S33.7. For example, the chloroformyl compound S33.6 is reacted with the amine S33.3 in a water-miscible solvent such as tetrahydrofuran, in the presence of aqueous sodium hydroxide, as described in *Org. Syn. Coll. Vol. 3*, 167, 1965, to yield the carbamate S33.7. Alternatively, the reaction is performed in dichloromethane in the presence of an organic base such as diisopropylethylamine or dimethylaminopyridine.

Scheme 33, Example 2 depicts the reaction of the chloroformate compound S33.6 with imidazole to produce the imidazolide S33.8. The imidazolide product is then reacted with the amine S33.3 to yield the carbamate S33.7. The preparation of the imidazolide is performed in an aprotic solvent such as dichloromethane at 0°, and the preparation of the carbamate is conducted in a similar solvent at ambient temperature, optionally in the presence of a base such as dimethylaminopyridine, as described in *J. Med. Chem.*, 1989, 32, 357.

Scheme 33 Example 3, depicts the reaction of the chloroformate S33.6 with an activated hydroxyl compound R"OH, to yield the mixed carbonate ester S33.10. The reaction is conducted in an inert organic solvent such as ether or dichloromethane, in the presence of a base such as dicyclohexylamine or triethylamine. The hydroxyl component R"OH is selected from the group of compounds S33.19 - S33.24 shown in Scheme 33, and similar compounds. For example, if the component R"OH is hydroxybenztriazole S33.19, N-hydroxysuccinimide S33.20, or pentachlorophenol, S33.21, the mixed carbonate S33.10 is obtained by the reaction of the chloroformate with the hydroxyl

compound in an ethereal solvent in the presence of dicyclohexylamine, as described in *Can. J. Chem.*, 1982, 60, 976. A similar reaction in which the component R"OH is pentafluorophenol S33.22 or 2-hydroxypyridine S33.23 is performed in an ethereal solvent in the presence of triethylamine, as described in
5 *Syn.*, 1986, 303, and *Chem. Ber.* 118, 468, 1985.

Scheme 33 Example 4 illustrates the preparation of carbamates in which an alkyloxycarbonylimidazole S33.8 is employed. In this procedure, an alcohol S33.5 is reacted with an equimolar amount of carbonyl diimidazole S33.11 to prepare the intermediate S33.8. The reaction is conducted in an aprotic organic
10 solvent such as dichloromethane or tetrahydrofuran. The acyloxyimidazole S33.8 is then reacted with an equimolar amount of the amine R'NH₂ to afford the carbamate S33.7. The reaction is performed in an aprotic organic solvent such as dichloromethane, as described in *Tet. Lett.*, 42, 2001, 5227, to afford the carbamate S33.7.

15 Scheme 33, Example 5 illustrates the preparation of carbamates by means of an intermediate alkoxycarbonylbenztriazole S33.13. In this procedure, an alcohol ROH is reacted at ambient temperature with an equimolar amount of benztriazole carbonyl chloride S33.12, to afford the alkoxycarbonyl product S33.13. The reaction is performed in an organic solvent such as benzene or
20 toluene, in the presence of a tertiary organic amine such as triethylamine, as described in *Synthesis.*, 1977, 704. The product is then reacted with the amine R'NH₂ to afford the carbamate S33.7. The reaction is conducted in toluene or ethanol, at from ambient temperature to about 80 °C as described in *Synthesis.*, 1977, 704.

25 Scheme 33, Example 6 illustrates the preparation of carbamates in which a carbonate (R"O)₂CO, S33.14, is reacted with an alcohol S33.5 to afford the intermediate alkyloxycarbonyl intermediate S33.15. The latter reagent is then reacted with the amine R'NH₂ to afford the carbamate S33.7. The procedure in which the reagent S33.15 is derived from hydroxybenztriazole S33.19 is
30 described in *Synthesis*, 1993, 908; the procedure in which the reagent S33.15 is derived from N-hydroxysuccinimide S33.20 is described in *Tet. Lett.*, 1992, 2781; the procedure in which the reagent S33.15 is derived from 2-hydroxypyridine S33.23 is described in *Tet. Lett.*, 1991, 4251; the procedure in

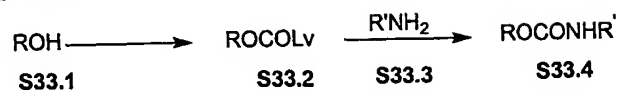
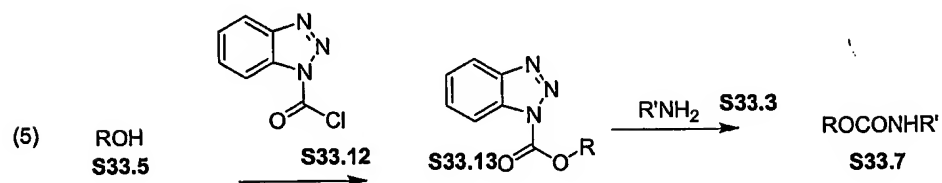
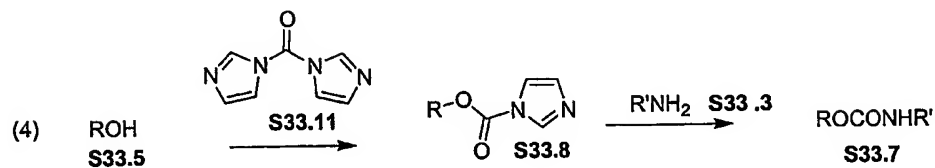
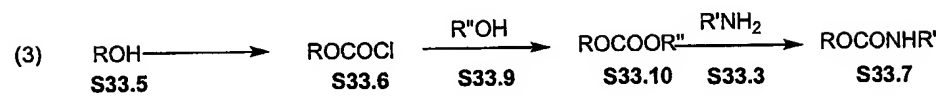
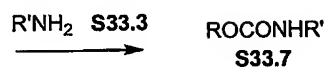
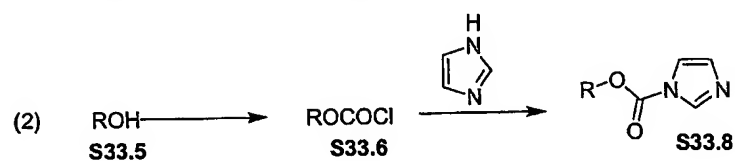
which the reagent S33.15 is derived from 4-nitrophenol S33.24 is described in *Synthesis*. 1993, 103. The reaction between equimolar amounts of the alcohol ROH and the carbonate S33.14 is conducted in an inert organic solvent at ambient temperature.

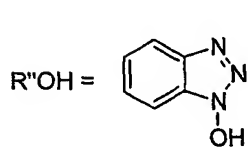
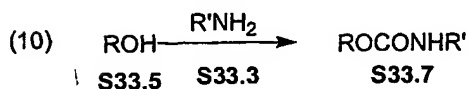
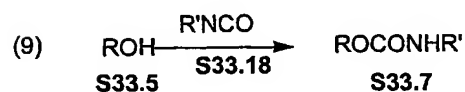
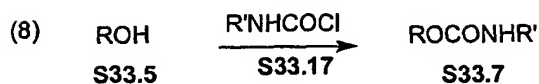
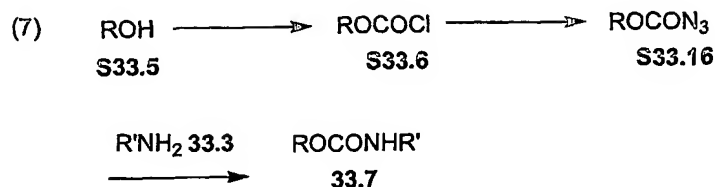
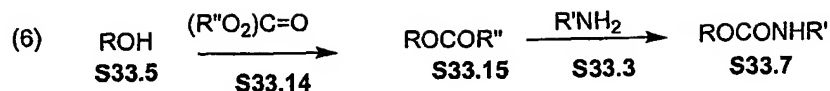
5 Scheme 33, Example 7 illustrates the preparation of carbamates from alkoxycarbonyl azides S33.16. In this procedure, an alkyl chloroformate S33.6 is reacted with an azide, for example sodium azide, to afford the alkoxycarbonyl azide S33.16. The latter compound is then reacted with an equimolar amount of the amine R'NH₂ to afford the carbamate S33.7. The reaction is conducted at
10 ambient temperature in a polar aprotic solvent such as dimethylsulfoxide, for example as described in *Synthesis*., 1982, 404.

 Scheme 33, Example 8 illustrates the preparation of carbamates by means of the reaction between an alcohol ROH and the chloroformyl derivative of an amine S33.17. In this procedure, which is described in Synthetic Organic
15 Chemistry, R. B. Wagner, H. D. Zook, Wiley, 1953, p. 647, the reactants are combined at ambient temperature in an aprotic solvent such as acetonitrile, in the presence of a base such as triethylamine, to afford the carbamate S33.7.

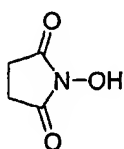
 Scheme 33, Example 9 illustrates the preparation of carbamates by means of the reaction between an alcohol ROH and an isocyanate S33.18. In this
20 procedure, which is described in Synthetic Organic Chemistry, R. B. Wagner, H. D. Zook, Wiley, 1953, p. 645, the reactants are combined at ambient temperature in an aprotic solvent such as ether or dichloromethane and the like, to afford the carbamate S33.7.

 Scheme 33, Example 10 illustrates the preparation of carbamates by
25 means of the reaction between an alcohol ROH and an amine R'NH₂. In this procedure, which is described in *Chem. Lett.* 1972, 373, the reactants are combined at ambient temperature in an aprotic organic solvent such as tetrahydrofuran, in the presence of a tertiary base such as triethylamine, and selenium. Carbon monoxide is passed through the solution and the reaction
30 proceeds to afford the carbamate S33.7.

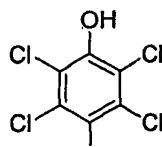
Scheme 33. Preparation of carbamates.**General reaction****Examples**



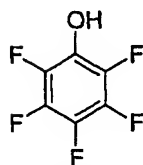
S33.19



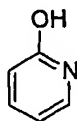
S33.20



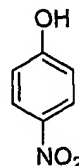
S33.21



S33.22



S33.23



S33.24

PREPARATION OF CARBOALKOXY-SUBSTITUTED PHOSPHONATE
BISAMIDATES, MONOAMIDATES, DIESTERS AND MONOESTERS.

- 5 A number of methods are available for the conversion of phosphonic acids into amidates and esters. In one group of methods, the phosphonic acid is either converted into an isolated activated intermediate such as a phosphoryl

chloride, or the phosphonic acid is activated in situ for reaction with an amine or a hydroxy compound.

The conversion of phosphonic acids into phosphoryl chlorides is accomplished by reaction with thionyl chloride, for example as described in *J. Gen. Chem. USSR*, 1983, 53, 480, *Zh. Obschei Khim.*, 1958, 28, 1063, or *J. Org. Chem.*, 1994, 59, 6144, or by reaction with oxalyl chloride, as described in *J. Am. Chem. Soc.*, 1994, 116, 3251, or *J. Org. Chem.*, 1994, 59, 6144, or by reaction with phosphorus pentachloride, as described in *J. Org. Chem.*, 2001, 66, 329, or in *J. Med. Chem.*, 1995, 38, 1372. The resultant phosphoryl chlorides are then reacted with amines or hydroxy compounds in the presence of a base to afford the amidate or ester products.

Phosphonic acids are converted into activated imidazolyl derivatives by reaction with carbonyl diimidazole, as described in *J. Chem. Soc., Chem. Comm.* (1991) 312, or *Nucleosides & Nucleotides* (2000) 19:1885. Activated sulfonyloxy derivatives are obtained by the reaction of phosphonic acids with trichloromethylsulfonyl chloride or with triisopropylbenzenesulfonyl chloride, as described in *Tet. Lett.* (1996) 7857, or *Bioorg. Med. Chem. Lett.* (1998) 8:663. The activated sulfonyloxy derivatives are then reacted with amines or hydroxy compounds to afford amidates or esters.

Alternatively, the phosphonic acid and the amine or hydroxy reactant are combined in the presence of a diimide coupling agent. The preparation of phosphonic amidates and esters by means of coupling reactions in the presence of dicyclohexyl carbodiimide is described, for example, in *J. Chem. Soc., Chem. Comm.* (1991) 312 or *Coll. Czech. Chem. Comm.* (1987) 52:2792. The use of ethyl dimethylaminopropyl carbodiimide for activation and coupling of phosphonic acids is described in *Tet. Lett.*, (2001) 42:8841, or *Nucleosides & Nucleotides* (2000) 19:1885.

A number of additional coupling reagents have been described for the preparation of amidates and esters from phosphonic acids. The agents include Aldrithiol-2, and PYBOP and BOP, as described in *J. Org. Chem.*, 1995, 60, 5214, and *J. Med. Chem.* (1997) 40:3842, mesitylene-2-sulfonyl-3-nitro-1,2,4-triazole (MSNT), as described in *J. Med. Chem.* (1996) 39:4958, diphenylphosphoryl azide, as described in *J. Org. Chem.* (1984) 49:1158, 1-

(2,4,6-triisopropylbenzenesulfonyl-3-nitro-1,2,4-triazole (TPSNT) as described in *Bioorg. Med. Chem. Lett.* (1998) 8:1013,

bromotris(dimethylamino)phosphonium hexafluorophosphate (BroP), as described in *Tet. Lett.*, (1996) 37:3997, 2-chloro-5,5-dimethyl-2-oxo-1,3,2-

5 dioxaphosphinane, as described in *Nucleosides Nucleotides* 1995, 14, 871, and diphenyl chlorophosphate, as described in *J. Med. Chem.*, 1988, 31, 1305.

Phosphonic acids are converted into amidates and esters by means of the Mitsunobu reaction, in which the phosphonic acid and the amine or hydroxy reactant are combined in the presence of a triaryl phosphine and a dialkyl
10 azodicarboxylate. The procedure is described in *Org. Lett.*, 2001, 3, 643, or *J. Med. Chem.*, 1997, 40, 3842.

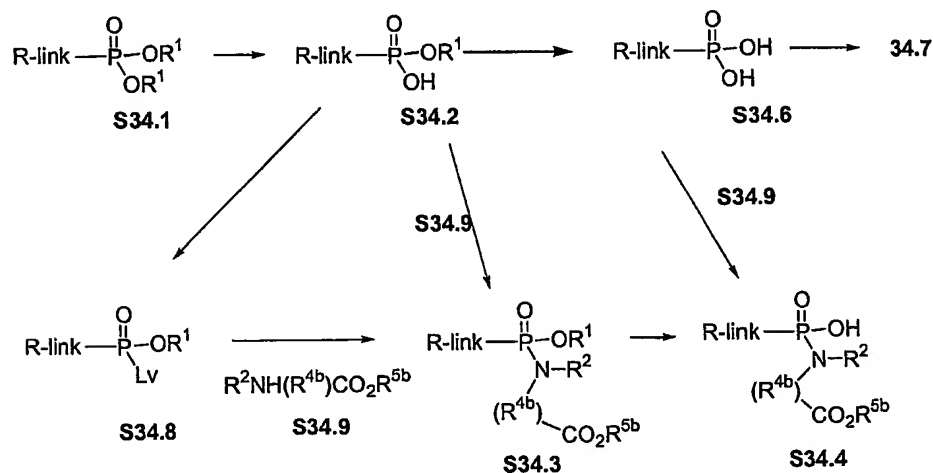
Phosphonic esters are also obtained by the reaction between phosphonic acids and halo compounds, in the presence of a suitable base. The method is described, for example, in *Anal. Chem.*, 1987, 59, 1056, or *J. Chem. Soc. Perkin
15 Trans., I*, 1993, 19, 2303, or *J. Med. Chem.*, 1995, 38, 1372, or *Tet. Lett.*, 2002, 43, 1161.

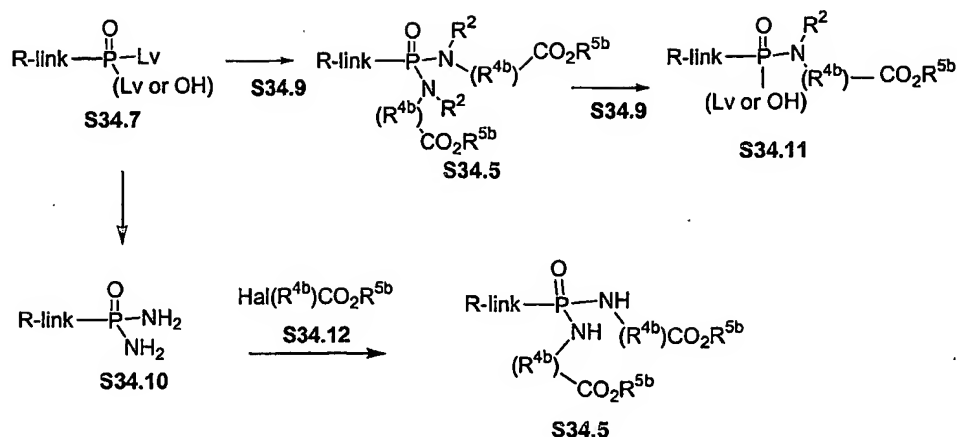
Schemes 34-37 illustrate the conversion of phosphonate esters and phosphonic acids into carboalkoxy-substituted phosphonbisamidates (Scheme 34), phosphonamidates (Scheme 35), phosphonate monoesters (Scheme 36) and
20 phosphonate diesters, (Scheme 37). Scheme 38 illustrates synthesis of gem-dialkyl amino phosphonate reagents.

Scheme 34 illustrates various methods for the conversion of phosphonate diesters S34.1 into phosphonbisamidates S34.5. The diester S34.1, prepared as described previously, is hydrolyzed, either to the monoester S34.2 or to the
25 phosphonic acid S34.6. The methods employed for these transformations are described above. The monoester S34.2 is converted into the monoamidate S34.3 by reaction with an aminoester S34.9, in which the group R² is H or alkyl; the group R^{4b} is a divalent alkylene moiety such as, for example, CHCH₃, CHCH₂CH₃, CH(CH(CH₃)₂), CH(CH₂Ph), and the like, or a side chain group
30 present in natural or modified aminoacids; and the group R^{5b} is C₁-C₁₂ alkyl, such as methyl, ethyl, propyl, isopropyl, or isobutyl; C₆-C₂₀ aryl, such as phenyl or substituted phenyl; or C₆-C₂₀ arylalkyl, such as benzyl or benzydryl. The reactants are combined in the presence of a coupling agent such as a

carbodiimide, for example dicyclohexyl carbodiimide, as described in *J. Am. Chem. Soc.*, (1957) 79:3575, optionally in the presence of an activating agent such as hydroxybenztriazole, to yield the amidate product S34.3. The amidate-forming reaction is also effected in the presence of coupling agents such as BOP, as described in *J. Org. Chem.* (1995) 60:5214, Aldrithiol, PYBOP and similar coupling agents used for the preparation of amides and esters. Alternatively, the reactants S34.2 and S34.9 are transformed into the monoamidate S34.3 by means of a Mitsunobu reaction. The preparation of amidates by means of the Mitsunobu reaction is described in *J. Med. Chem.* (1995) 38:2742. Equimolar amounts of the reactants are combined in an inert solvent such as tetrahydrofuran in the presence of a triaryl phosphine and a dialkyl azodicarboxylate. The thus-obtained monoamidate ester S34.3 is then transformed into amidate phosphonic acid S34.4. The conditions used for the hydrolysis reaction depend on the nature of the R¹ group, as described previously. The phosphonic acid amidate S34.4 is then reacted with an aminoester S34.9, as described above, to yield the bisamidate product S34.5, in which the amino substituents are the same or different. Alternatively, the phosphonic acid S34.6 may be treated with two different amino ester reagents simultaneously, i.e. S34.9 where R², R^{4b} or R^{5b} are different. The resulting mixture of bisamidate products S34.5 may then be separable, e.g. by chromatography.

Scheme 34





An example of this procedure is shown in Scheme 34, Example 1. In this procedure, a dibenzyl phosphonate **S34.14** is reacted with diazabicyclooctane (DABCO) in toluene at reflux, as described in *J. Org. Chem.*, 1995, 60, 2946, to afford the monobenzyl phosphonate **S34.15**. The product is then reacted with equimolar amounts of ethyl alaninate **S34.16** and dicyclohexyl carbodiimide in pyridine, to yield the amidate product **S34.17**. The benzyl group is then removed, for example by hydrogenolysis over a palladium catalyst, to give the monoacid product **S34.18** which may be unstable according to *J. Med. Chem.* (1997) 40(23):3842. This compound **S34.18** is then reacted in a Mitsunobu reaction with ethyl leucinate **S34.19**, triphenyl phosphine and diethylazodicarboxylate, as described in *J. Med. Chem.*, 1995, 38, 2742, to produce the bisamidate product **S34.20**.

Using the above procedures, but employing in place of ethyl leucinate **S34.19** or ethyl alaninate **S34.16**, different aminoesters **S34.9**, the corresponding products **S34.5** are obtained.

Alternatively, the phosphonic acid **S34.6** is converted into the bisamidate **S34.5** by use of the coupling reactions described above. The reaction is performed in one step, in which case the nitrogen-related substituents present in the product **S34.5** are the same, or in two steps, in which case the nitrogen-related substituents can be different.

An example of the method is shown in Scheme 34, Example 2. In this procedure, a phosphonic acid **S34.6** is reacted in pyridine solution with excess ethyl phenylalaninate **S34.21** and dicyclohexylcarbodiimide, for example as

described in *J. Chem. Soc., Chem. Comm.*, 1991, 1063, to give the bisamidate product S34.22.

Using the above procedures, but employing, in place of ethyl phenylalaninate, different aminoesters S34.9, the corresponding products S34.5 are obtained.

As a further alternative, the phosphonic acid S34.6 is converted into the mono or bis-activated derivative S34.7, in which Lv is a leaving group such as chloro, imidazolyl, triisopropylbenzenesulfonyloxy etc. The conversion of phosphonic acids into chlorides S34.7 (Lv = Cl) is effected by reaction with thionyl chloride or oxalyl chloride and the like, as described in Organic Phosphorus Compounds, G. M. Kosolapoff, L. Maeir, eds, Wiley, 1976, p. 17. The conversion of phosphonic acids into monoimidazolides S34.7 (Lv = imidazolyl) is described in *J. Med. Chem.*, 2002, 45, 1284 and in *J. Chem. Soc. Chem. Comm.*, 1991, 312. Alternatively, the phosphonic acid is activated by reaction with triisopropylbenzenesulfonyl chloride, as described in *Nucleosides and Nucleotides*, 2000, 10, 1885. The activated product is then reacted with the aminoester S34.9, in the presence of a base, to give the bisamidate S34.5. The reaction is performed in one step, in which case the nitrogen substituents present in the product S34.5 are the same, or in two steps, via the intermediate S34.11, in which case the nitrogen substituents can be different.

Examples of these methods are shown in Scheme 34, Examples 3 and 5. In the procedure illustrated in Scheme 34, Example 3, a phosphonic acid S34.6 is reacted with ten molar equivalents of thionyl chloride, as described in *Zh. Obschei Khim.*, 1958, 28, 1063, to give the dichloro compound S34.23. The product is then reacted at reflux temperature in a polar aprotic solvent such as acetonitrile, and in the presence of a base such as triethylamine, with butyl serinate S34.24 to afford the bisamidate product S34.25.

Using the above procedures, but employing, in place of butyl serinate S34.24, different aminoesters S34.9, the corresponding products S34.5 are obtained.

In the procedure illustrated in Scheme 34, Example 5, the phosphonic acid S34.6 is reacted, as described in *J. Chem. Soc. Chem. Comm.*, 1991, 312, with carbonyl diimidazole to give the imidazolid S34.S32. The product is then

reacted in acetonitrile solution at ambient temperature, with one molar equivalent of ethyl alaninate S34.33 to yield the monodisplacement product S34.34. The latter compound is then reacted with carbonyl diimidazole to produce the activated intermediate S34.35, and the product is then reacted, under
5 the same conditions, with ethyl N-methylalaninate S34.33a to give the bisamidate product S34.36.

Using the above procedures, but employing, in place of ethyl alaninate S34.33 or ethyl N-methylalaninate S34.33a, different aminoesters S34.9, the corresponding products S34.5 are obtained.

10 The intermediate monoamidate S34.3 is also prepared from the monoester S34.2 by first converting the monoester into the activated derivative S34.8 in which Lv is a leaving group such as halo, imidazolyl etc, using the procedures described above. The product S34.8 is then reacted with an aminoester S34.9 in the presence of a base such as pyridine, to give an
15 intermediate monoamidate product S34.3. The latter compound is then converted, by removal of the R¹ group and coupling of the product with the aminoester S34.9, as described above, into the bisamidate S34.5.

An example of this procedure, in which the phosphonic acid is activated by conversion to the chloro derivative S34.26, is shown in Scheme 34, Example
20 4. In this procedure, the phosphonic monobenzyl ester S34.15 is reacted, in dichloromethane, with thionyl chloride, as described in *Tet. Letters.*, 1994, 35, 4097, to afford the phosphoryl chloride S34.26. The product is then reacted in acetonitrile solution at ambient temperature with one molar equivalent of ethyl 3-amino-2-methylpropionate S34.27 to yield the monoamidate product S34.28.
25 The latter compound is hydrogenated in ethylacetate over a 5% palladium on carbon catalyst to produce the monoacid product S34.29. The product is subjected to a Mitsunobu coupling procedure, with equimolar amounts of butyl alaninate S34.30, triphenyl phosphine, diethylazodicarboxylate and triethylamine in tetrahydrofuran, to give the bisamidate product S34.31.

30 Using the above procedures, but employing, in place of ethyl 3-amino-2-methylpropionate S34.27 or butyl alaninate S34.30, different aminoesters S34.9, the corresponding products S34.5 are obtained.

The activated phosphonic acid derivative S34.7 is also converted into the bisamidate S34.5 via the diamino compound S34.10. The conversion of activated phosphonic acid derivatives such as phosphoryl chlorides into the corresponding amino analogs S34.10, by reaction with ammonia, is described in
5 Organic Phosphorus Compounds, G. M. Kosolapoff, L. Maeir, eds, Wiley, 1976. The bisamino compound S34.10 is then reacted at elevated temperature with a haloester S34.12 (Hal = halogen, i.e. F, Cl, Br, I), in a polar organic solvent such as dimethylformamide, in the presence of a base such as 4, 4-dimethylaminopyridine (DMAP) or potassium carbonate, to yield the bisamidate
10 S34.5. Alternatively, S34.6 may be treated with two different amino ester reagents simulataneously, i.e. S34.12 where R^{4b} or R^{5b} are different. The resulting mixture of bisamidate products S34.5 may then be separable, e.g. by chromatography.

An example of this procedure is shown in Scheme 34, Example 6. In this
15 method, a dichlorophosphonate S34.23 is reacted with ammonia to afford the diamide S34.37. The reaction is performed in aqueous, aqueous alcoholic or alcoholic solution, at reflux temperature. The resulting diamino compound is then reacted with two molar equivalents of ethyl 2-bromo-3-methylbutyrate S34.38, in a polar organic solvent such as N-methylpyrrolidinone at ca. 150 °C,
20 in the presence of a base such as potassium carbonate, and optionally in the presence of a catalytic amount of potassium iodide, to afford the bisamidate product S34.39.

Using the above procedures, but employing, in place of ethyl 2-bromo-3-methylbutyrate S34.38, different haloesters S34.12 the corresponding products
25 S34.5 are obtained.

The procedures shown in Scheme 34 are also applicable to the preparation of bisamidates in which the aminoester moiety incorporates different functional groups. Scheme 34, Example 7 illustrates the preparation of bisamidates derived from tyrosine. In this procedure, the monoimidazolidine
30 S34.32 is reacted with propyl tyrosinate S34.40, as described in Example 5, to yield the monoamidate S34.41. The product is reacted with carbonyl diimidazole to give the imidazolidine S34.42, and this material is reacted with a further molar equivalent of propyl tyrosinate to produce the bisamidate product S34.43.

Using the above procedures, but employing, in place of propyl tyrosinate S34.40, different aminoesters S34.9, the corresponding products S34.5 are obtained. The aminoesters employed in the two stages of the above procedure can be the same or different, so that bisamidates with the same or different amino substituents are prepared.

Scheme 35 illustrates methods for the preparation of phosphonate monoamidates.

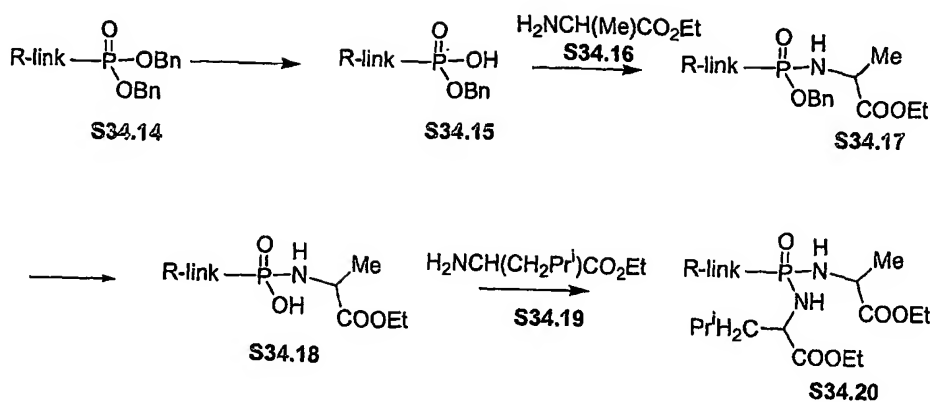
In one procedure, a phosphonate monoester S34.1 is converted, as described in Scheme 34, into the activated derivative S34.8. This compound is then reacted, as described above, with an aminoester S34.9, in the presence of a base, to afford the monoamidate product S35.1.

The procedure is illustrated in Scheme 35, Example 1. In this method, a monophenyl phosphonate S35.7 is reacted with, for example, thionyl chloride, as described in *J. Gen. Chem. USSR.*, 1983, 32, 367, to give the chloro product S35.8. The product is then reacted, as described in Scheme 34, with ethyl alaninate, S3, to yield the amidate S35.10.

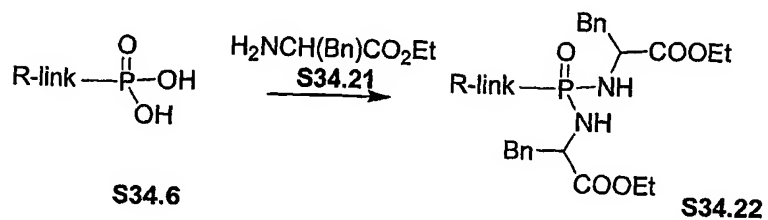
Using the above procedures, but employing, in place of ethyl alaninate S35.9, different aminoesters S34.9, the corresponding products S35.1 are obtained.

Alternatively, the phosphonate monoester S34.1 is coupled, as described in Scheme 34, with an aminoester S34.9 to produce the amidate, S335.1. If necessary, the R¹ substituent is then altered, by initial cleavage to afford the phosphonic acid S35.2. The procedures for this transformation depend on the nature of the R¹ group, and are described above. The phosphonic acid is then transformed into the ester amidate product S35.3, by reaction with the hydroxy compound R³OH, in which the group R³ is aryl, heterocycle, alkyl, cycloalkyl, haloalkyl etc, using the same coupling procedures (carbodiimide, Aldrithiol-2, PYBOP, Mitsunobu reaction etc) described in Scheme 34 for the coupling of amines and phosphonic acids.

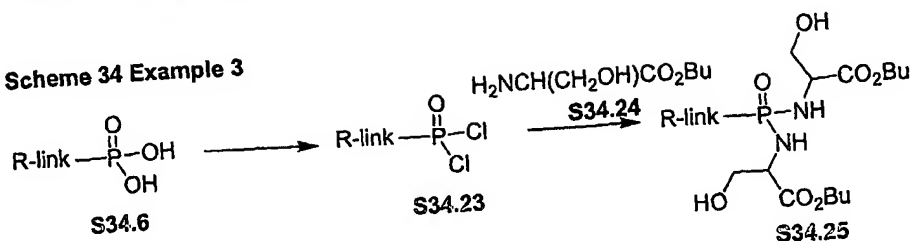
Scheme 34 Example 1



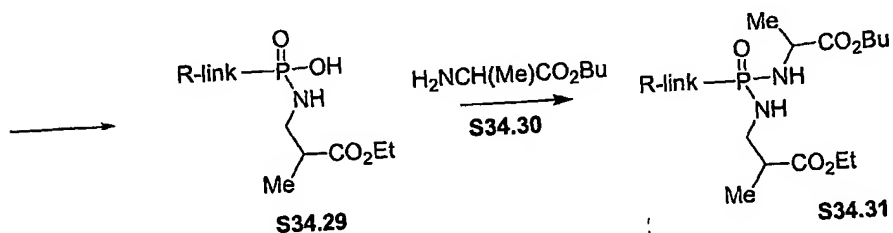
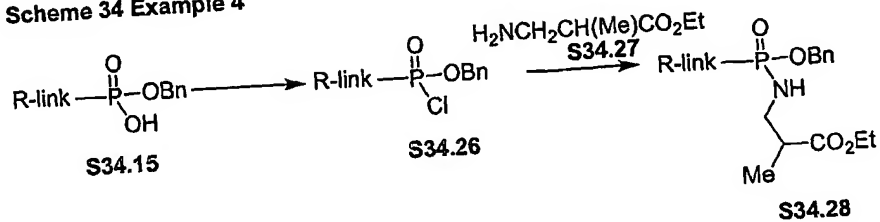
Scheme 34 Example 2



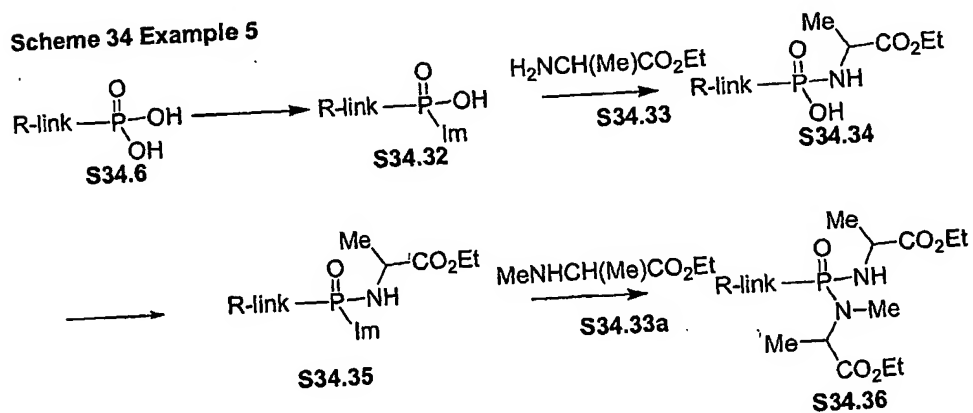
Scheme 34 Example 3



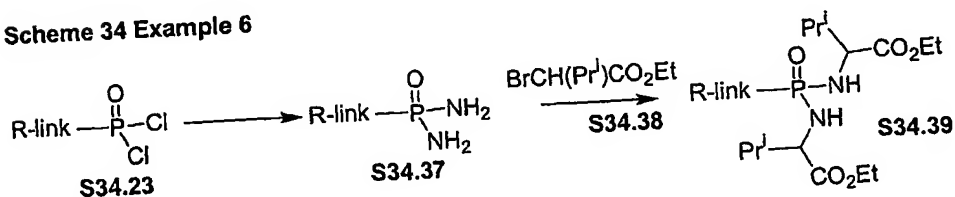
Scheme 34 Example 4



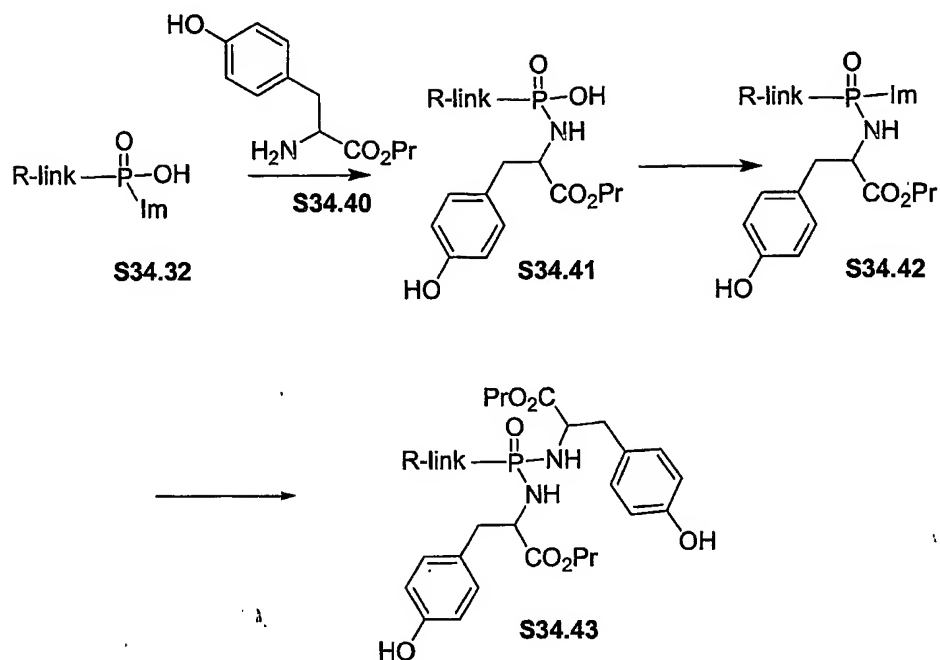
Scheme 34 Example 5



Scheme 34 Example 6



Scheme 34 Example 7



Examples of this method are shown in Scheme 35, Examples and 2 and
 5 3. In the sequence shown in Example 2, a monobenzyl phosphonate S35.11 is
 transformed by reaction with ethyl alaninate, using one of the methods described
 above, into the monoamidate S35.12. The benzyl group is then removed by
 catalytic hydrogenation in ethylacetate solution over a 5% palladium on carbon
 catalyst, to afford the phosphonic acid amidate S35.13. The product is then
 10 reacted in dichloromethane solution at ambient temperature with equimolar
 amounts of 1-(dimethylaminopropyl)-3-ethylcarbodiimide and trifluoroethanol
 S35.14, for example as described in *Tet. Lett.*, 2001, 42, 8841, to yield the
 amidate ester S35.15.

In the sequence shown in Scheme 35, Example 3, the monoamidate
 15 S35.13 is coupled, in tetrahydrofuran solution at ambient temperature, with
 equimolar amounts of dicyclohexyl carbodiimide and 4-hydroxy-N-
 methylpiperidine S35.16, to produce the amidate ester product S35.17.

Using the above procedures, but employing, in place of the ethyl
 alaninate product S35.12 different monoacids S35.2, and in place of

trifluoroethanol S35.14 or 4-hydroxy-N-methylpiperidine S35.16, different hydroxy compounds R^3OH , the corresponding products S35.3 are obtained.

Alternatively, the activated phosphonate ester S34.8 is reacted with ammonia to yield the amidate S35.4. The product is then reacted, as described in
5 Scheme 34, with a haloester S35.5, in the presence of a base, to produce the amidate product S35.6. If appropriate, the nature of the R^1 group is changed, using the procedures described above, to give the product S35.3. The method is illustrated in Scheme 35, Example 4. In this sequence, the monophenyl phosphoryl chloride S35.18 is reacted, as described in Scheme 34, with
10 ammonia, to yield the amino product S35.19. This material is then reacted in N-methylpyrrolidinone solution at 170° with butyl 2-bromo-3-phenylpropionate S35.20 and potassium carbonate, to afford the amidate product S35.21.

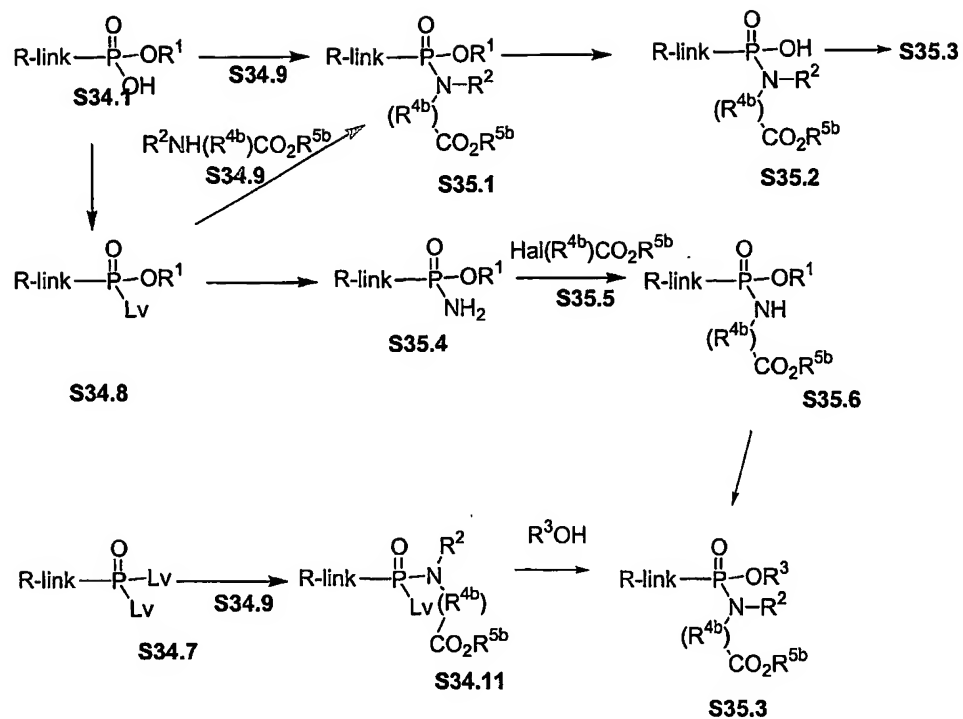
Using these procedures, but employing, in place of butyl 2-bromo-3-phenylpropionate S35.20, different haloesters S35.5, the corresponding products
15 S35.6 are obtained.

The monoamidate products S35.3 are also prepared from the doubly activated phosphonate derivatives S34.7. In this procedure, examples of which are described in *Synlett.*, 1998, 1, 73, the intermediate S34.7 is reacted with a limited amount of the aminoester S34.9 to give the mono-displacement product
20 S34.11. The latter compound is then reacted with the hydroxy compound R^3OH in a polar organic solvent such as dimethylformamide, in the presence of a base such as diisopropylethylamine, to yield the monoamidate ester S35.3.

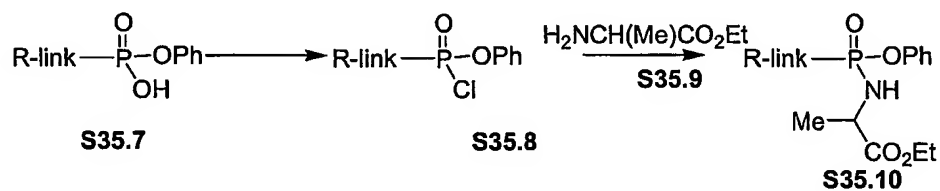
The method is illustrated in Scheme 35, Example 5. In this method, the phosphoryl dichloride S35.22 is reacted in dichloromethane solution with one
25 molar equivalent of ethyl N-methyl tyrosinate S35.23 and dimethylaminopyridine, to generate the monoamidate S35.24. The product is then reacted with phenol S35.25 in dimethylformamide containing potassium carbonate, to yield the ester amidate product S35.26.

Using these procedures, but employing, in place of ethyl N-methyl tyrosinate S35.23 or phenol S35.25, the aminoesters 34.9 and/or the hydroxy
30 compounds R^3OH , the corresponding products S35.3 are obtained.

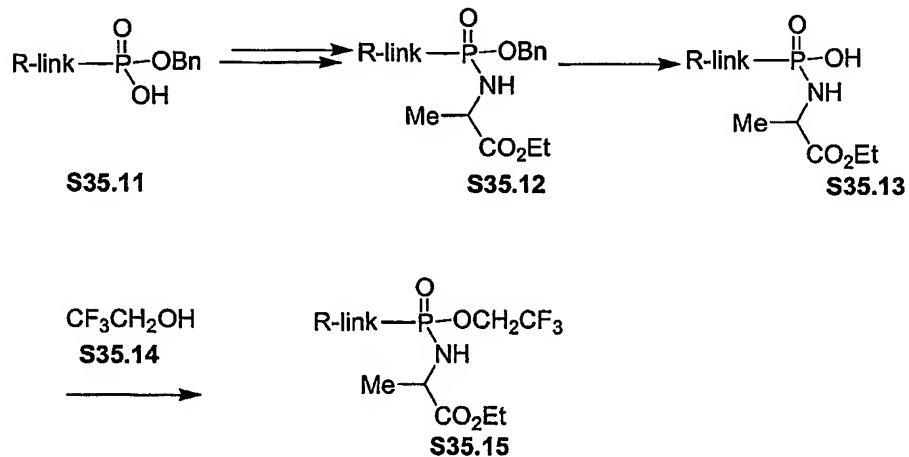
Scheme 35



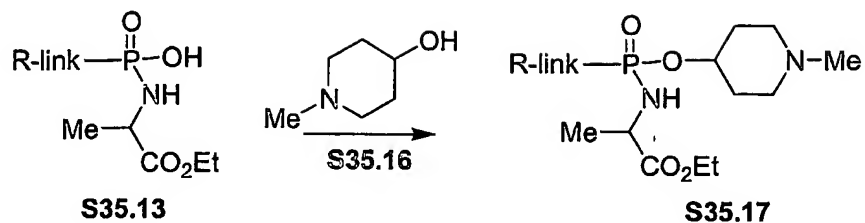
Scheme 35 Example 1



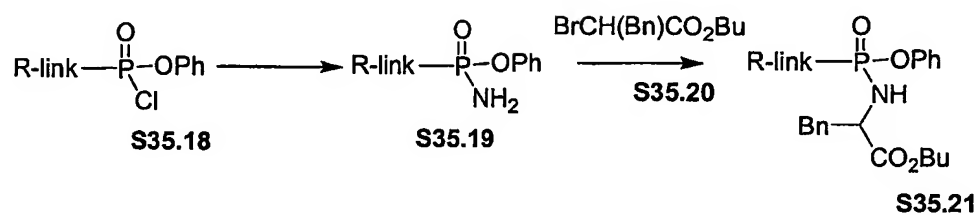
Scheme 35 Example 2



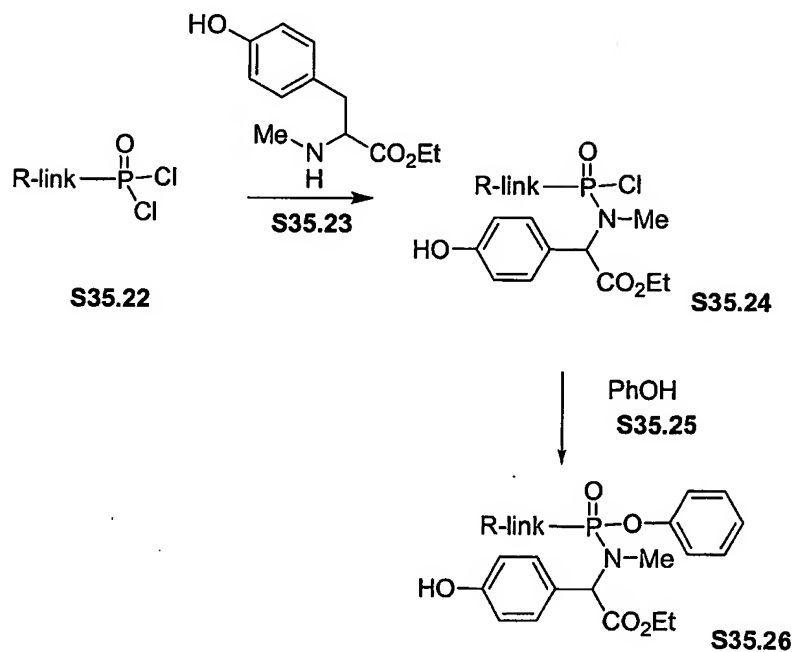
Scheme 35 Example 3



Scheme 35 Example 4



Scheme 35 Example 5



5

Scheme 36 illustrates methods for the preparation of carboalkoxy-substituted phosphonate diesters in which one of the ester groups incorporates a carboalkoxy substituent.

In one procedure, a phosphonate monoester S34.1, prepared as described above, is coupled, using one of the methods described above, with a hydroxyester S36.1, in which the groups R^{4b} and R^{5b} are as described in Scheme 34. For example, equimolar amounts of the reactants are coupled in the presence of a carbodiimide such as dicyclohexyl carbodiimide, as described in *Aust. J. Chem.*, 1963, 609, optionally in the presence of dimethylaminopyridine, as described in *Tet.*, 1999, 55, 12997. The reaction is conducted in an inert solvent at ambient temperature.

The procedure is illustrated in Scheme 36, Example 1. In this method, a monophenyl phosphonate S36.9 is coupled, in dichloromethane solution in the presence of dicyclohexyl carbodiimide, with ethyl 3-hydroxy-2-methylpropionate S36.10 to yield the phosphonate mixed diester S36.11.

Using this procedure, but employing, in place of ethyl 3-hydroxy-2-methylpropionate S36.10, different hydroxyesters S33.1, the corresponding products S33.2 are obtained.

The conversion of a phosphonate monoester S34.1 into a mixed diester S36.2 is also accomplished by means of a Mitsunobu coupling reaction with the hydroxyester S36.1, as described in *Org. Lett.*, 2001, 643. In this method, the reactants S34.1 and S36.1 are combined in a polar solvent such as tetrahydrofuran, in the presence of a triarylphosphine and a dialkyl azodicarboxylate, to give the mixed diester S36.2. The R¹ substituent is varied by cleavage, using the methods described previously, to afford the monoacid product S36.3. The product is then coupled, for example using methods described above, with the hydroxy compound R³OH, to give the diester product S36.4.

The procedure is illustrated in Scheme 36, Example 2. In this method, a monoallyl phosphonate S36.12 is coupled in tetrahydrofuran solution, in the presence of triphenylphosphine and diethylazodicarboxylate, with ethyl lactate S36.13 to give the mixed diester S36.14. The product is reacted with tris(triphenylphosphine) rhodium chloride (Wilkinson catalyst) in acetonitrile, as described previously, to remove the allyl group and produce the monoacid product S36.15. The latter compound is then coupled, in pyridine solution at ambient temperature, in the presence of dicyclohexyl carbodiimide, with one molar equivalent of 3-hydroxypyridine S36.16 to yield the mixed diester S36.17.

Using the above procedures, but employing, in place of the ethyl lactate S36.13 or 3-hydroxypyridine, a different hydroxyester S36.1 and/or a different hydroxy compound R³OH, the corresponding products S36.4 are obtained.

The mixed diesters S36.2 are also obtained from the monoesters S34.1
5 via the intermediacy of the activated monoesters S36.5. In this procedure, the monoester S34.1 is converted into the activated compound S36.5 by reaction with, for example, phosphorus pentachloride, as described in *J. Org. Chem.*, 2001, 66, 329, or with thionyl chloride or oxalyl chloride (Lv = Cl), or with triisopropylbenzenesulfonyl chloride in pyridine, as described in *Nucleosides and Nucleotides*, 2000, 19, 1885, or with carbonyl diimidazole, as described in *J.*
10 *Med. Chem.*, 2002, 45, 1284. The resultant activated monoester is then reacted with the hydroxyester S36.1, as described above, to yield the mixed diester S36.2.

The procedure is illustrated in Scheme 36, Example 3. In this sequence, a
15 monophenyl phosphonate S36.9 is reacted, in acetonitrile solution at 70 °C, with ten equivalents of thionyl chloride, so as to produce the phosphoryl chloride S36.19. The product is then reacted with ethyl 4-carbamoyl-2-hydroxybutyrate S36.20 in dichloromethane containing triethylamine, to give the mixed diester S36.21.

20 Using the above procedures, but employing, in place of ethyl 4-carbamoyl-2-hydroxybutyrate S36.20, different hydroxyesters S36.1, the corresponding products S36.2 are obtained.

The mixed phosphonate diesters are also obtained by an alternative route for incorporation of the R³O group into intermediates S36.3 in which the
25 hydroxyester moiety is already incorporated. In this procedure, the monoacid intermediate S36.3 is converted into the activated derivative S36.6 in which Lv is a leaving group such as chloro, imidazole, and the like, as previously described. The activated intermediate is then reacted with the hydroxy compound R³OH, in the presence of a base, to yield the mixed diester product
30 S36.4.

The method is illustrated in Scheme 36, Example 4. In this sequence, the phosphonate monoacid S36.22 is reacted with trichloromethanesulfonyl chloride in tetrahydrofuran containing collidine, as described in *J. Med. Chem.*, 1995, 38,

4648, to produce the trichloromethanesulfonyloxy product **S36.23**. This compound is reacted with 3-(morpholinomethyl)phenol **S36.24** in dichloromethane containing triethylamine, to yield the mixed diester product **S36.25**.

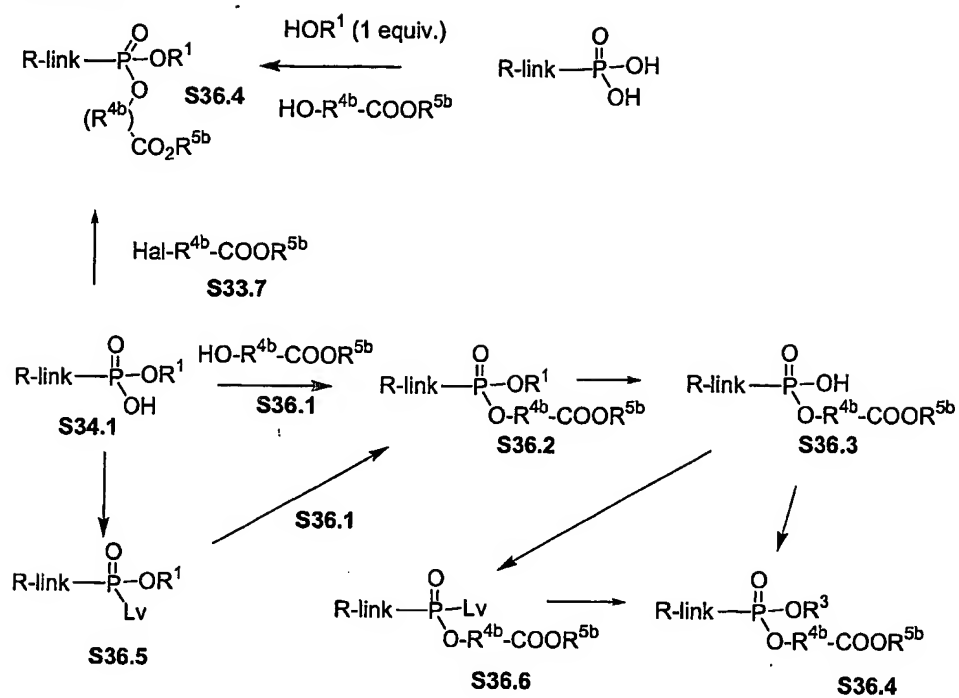
- 5 Using the above procedures, but employing, in place of with 3-(morpholinomethyl)phenol **S36.24**, different alcohols R^3OH , the corresponding products **S36.4** are obtained.

 The phosphonate esters **S36.4** are also obtained by means of alkylation reactions performed on the monoesters **S34.1**. The reaction between the
10 monoacid **S34.1** and the haloester **S36.7** is performed in a polar solvent in the presence of a base such as diisopropylethylamine, as described in *Anal. Chem.*, 1987, 59, 1056, or triethylamine, as described in *J. Med. Chem.*, 1995, 38, 1372, or in a non-polar solvent such as benzene, in the presence of 18-crown-6, as described in *Syn. Comm.*, 1995, 25, 3565.

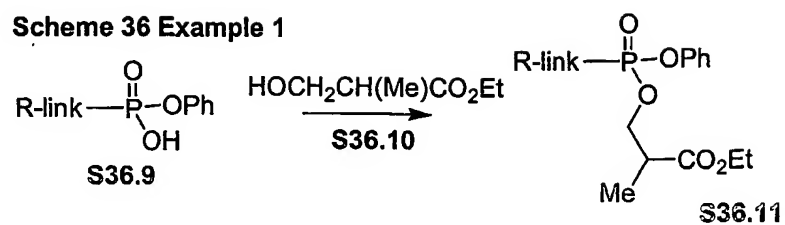
- 15 The method is illustrated in Scheme 36, Example 5. In this procedure, the monoacid **S36.26** is reacted with ethyl 2-bromo-3-phenylpropionate **S36.27** and diisopropylethylamine in dimethylformamide at 80 °C to afford the mixed diester product **S36.28**.

 Using the above procedure, but employing, in place of ethyl 2-bromo-3-
20 phenylpropionate **S36.27**, different haloesters **S36.7**, the corresponding products **S36.4** are obtained.

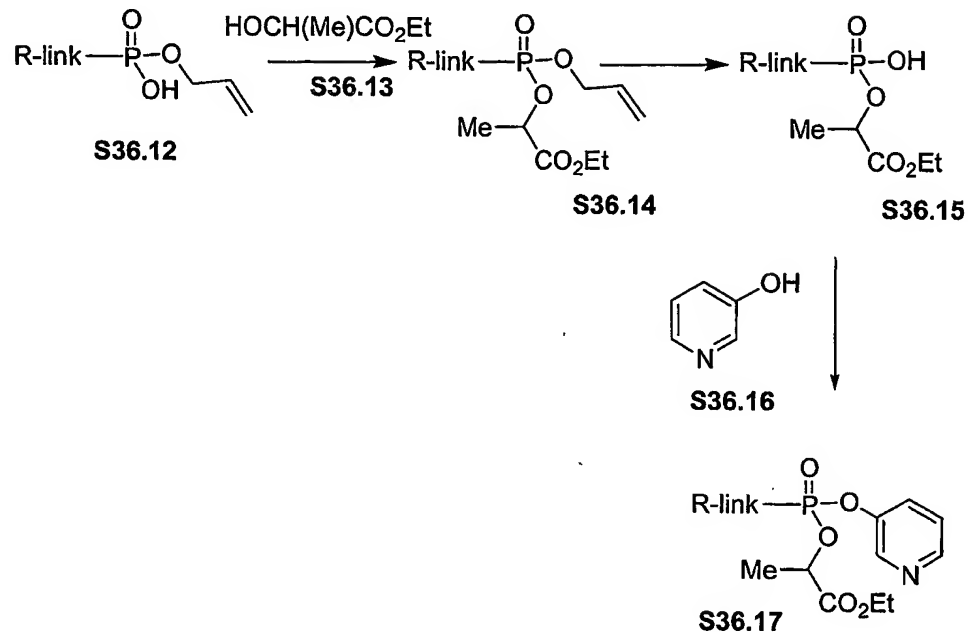
Scheme 36



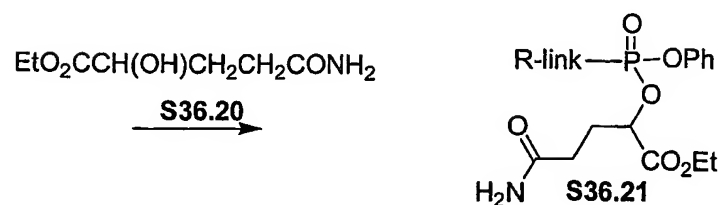
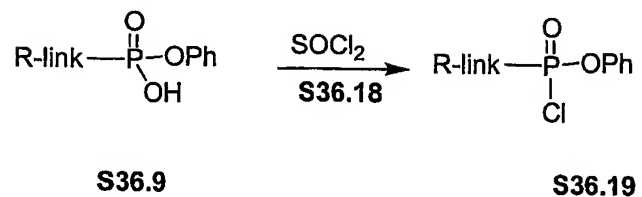
Scheme 36 Example 1



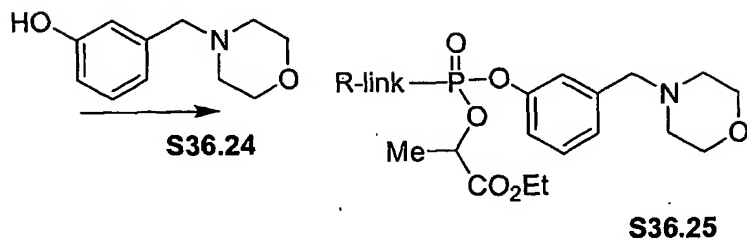
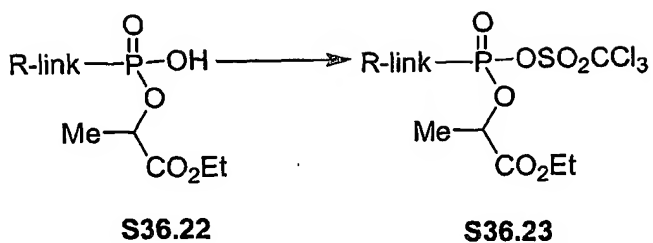
Scheme 36 Example 2



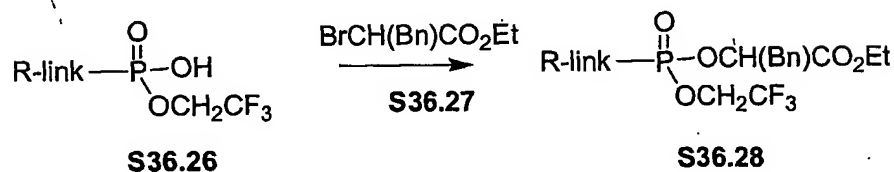
Scheme 36 Example 3



Scheme 36 Example 4



Scheme 36 Example 5



Scheme 37 illustrates methods for the preparation of phosphonate diesters in which both the ester substituents incorporate carboalkoxy groups.

The compounds are prepared directly or indirectly from the phosphonic acids **S34.6**. In one alternative, the phosphonic acid is coupled with the hydroxyester **S37.2**, using the conditions described previously in Schemes 34-36, such as coupling reactions using dicyclohexyl carbodiimide or similar reagents, or under the conditions of the Mitsunobu reaction, to afford the diester product **S37.3** in which the ester substituents are identical.

This method is illustrated in Scheme 37, Example 1. In this procedure, the phosphonic acid **S34.6** is reacted with three molar equivalents of butyl lactate

S37.5 in the presence of Aldrithiol-2 and triphenyl phosphine in pyridine at ca. 70 °C, to afford the diester S37.6.

Using the above procedure, but employing, in place of butyl lactate S37.5, different hydroxyesters S37.2, the corresponding products S37.3 are
5 obtained.

Alternatively, the diesters S37.3 are obtained by alkylation of the phosphonic acid S34.6 with a haloester S37.1. The alkylation reaction is performed as described in Scheme 36 for the preparation of the esters S36.4.

This method is illustrated in Scheme 37, Example 2. In this procedure,
10 the phosphonic acid S34.6 is reacted with excess ethyl 3-bromo-2-methylpropionate S37.7 and diisopropylethylamine in dimethylformamide at ca. 80 °C, as described in *Anal. Chem.*, 1987, 59, 1056, to produce the diester S37.8.

Using the above procedure, but employing, in place of ethyl 3-bromo-2-methylpropionate S37.7, different haloesters S37.1, the corresponding products
15 S37.3 are obtained.

The diesters S37.3 are also obtained by displacement reactions of activated derivatives S34.7 of the phosphonic acid with the hydroxyesters S37.2. The displacement reaction is performed in a polar solvent in the presence of a suitable base, as described in Scheme 36. The displacement reaction is
20 performed in the presence of an excess of the hydroxyester, to afford the diester product S37.3 in which the ester substituents are identical, or sequentially with limited amounts of different hydroxyesters, to prepare diesters S37.3 in which the ester substituents are different.

The methods are illustrated in Scheme 37, Examples 3 and 4. As shown
25 in Example 3, the phosphoryl dichloride S35.22 is reacted with three molar equivalents of ethyl 3-hydroxy-2-(hydroxymethyl)propionate S37.9 in tetrahydrofuran containing potassium carbonate, to obtain the diester product S37.10.

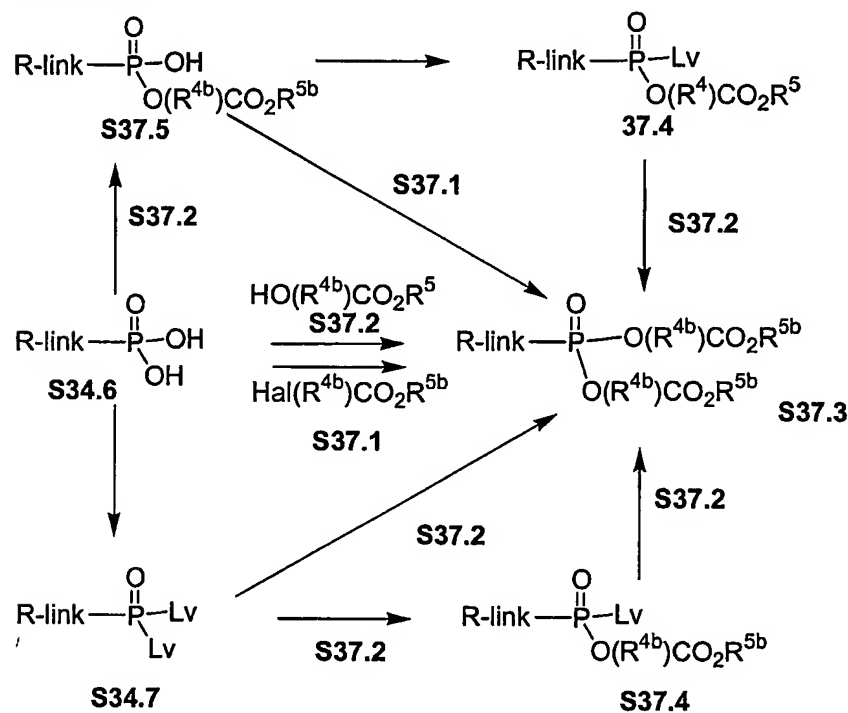
Using the above procedure, but employing, in place of ethyl 3-hydroxy-
30 2-(hydroxymethyl)propionate S37.9, different hydroxyesters S37.2, the corresponding products S37.3 are obtained.

Scheme 37, Example 4 depicts the displacement reaction between equimolar amounts of the phosphoryl dichloride S35.22 and ethyl 2-methyl-3-

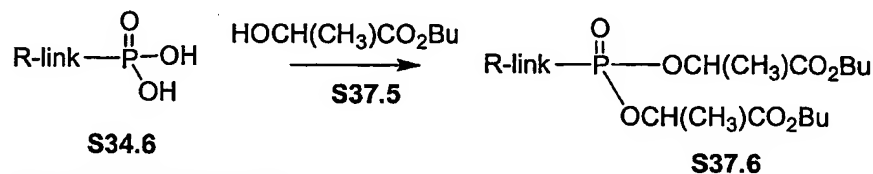
hydroxypropionate S37.11, to yield the monoester product S37.12. The reaction is conducted in acetonitrile at 70° in the presence of diisopropylethylamine. The product S37.12 is then reacted, under the same conditions, with one molar equivalent of ethyl lactate S37.13, to give the diester product S37.14.

- 5 Using the above procedures, but employing, in place of ethyl 2-methyl-3-hydroxypropionate S37.11 and ethyl lactate S37.13, sequential reactions with different hydroxyesters S37.2, the corresponding products S37.3 are obtained.

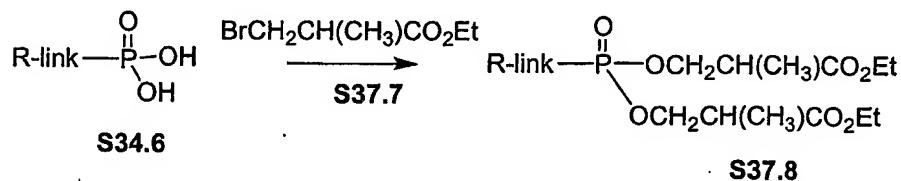
Scheme 37



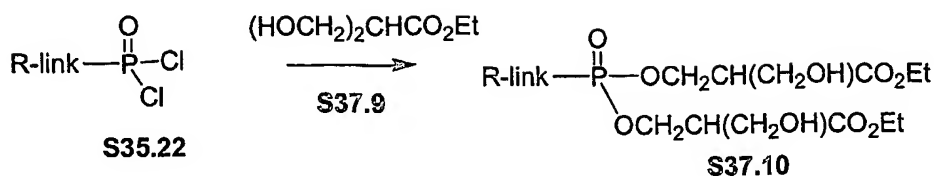
Scheme 37 Example 1



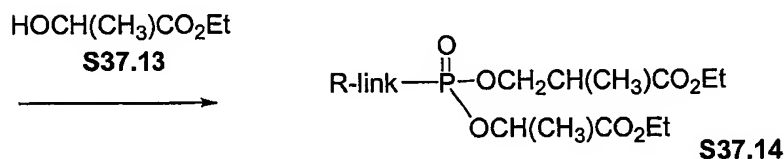
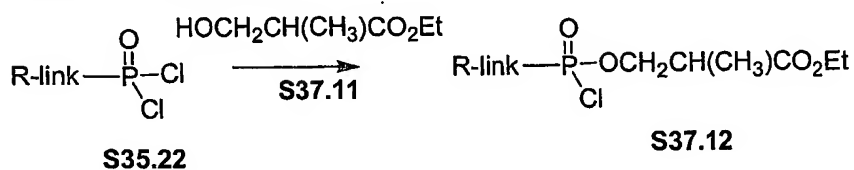
Scheme 37 Example 2



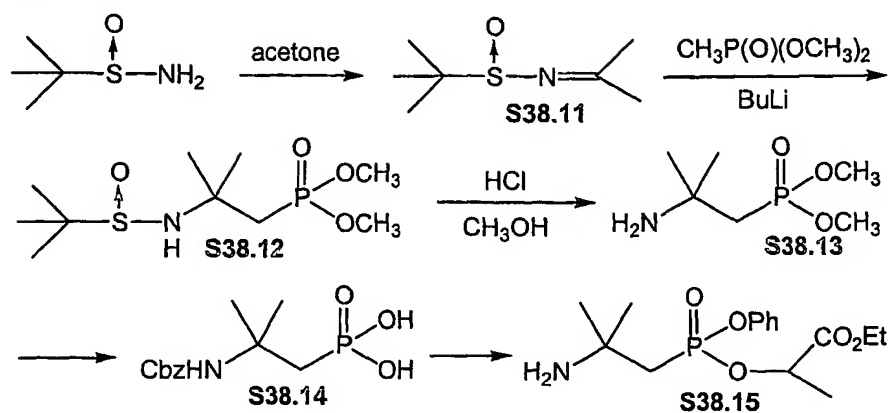
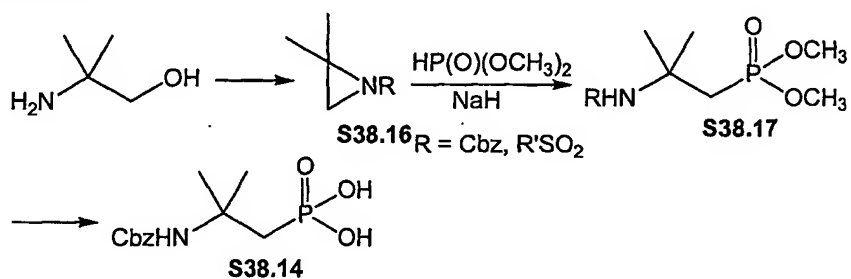
Scheme 37 Example 3



Scheme 37 Example 4

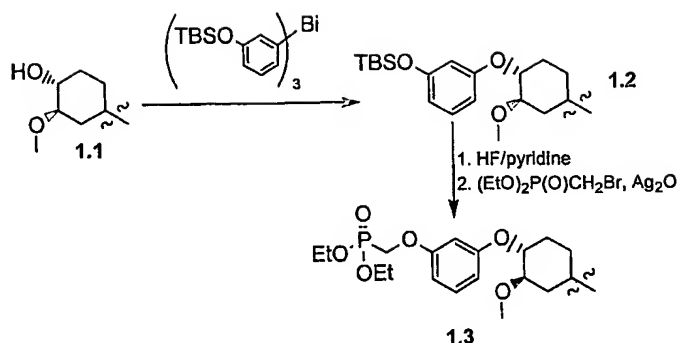


- 2,2-Dimethyl-2-aminoethylphosphonic acid intermediates can be prepared by the route in Scheme 5. Condensation of 2-methyl-2-propanesulfonamide with acetone give sulfinyl imine S38.11 (*J. Org. Chem.* 1999, 64, 12). Addition of dimethyl methylphosphonate lithium to S38.11 afford S38.12. Acidic methanolysis of S38.12 provide amine S38.13. Protection of amine with Cbz group and removal of methyl groups yield phosphonic acid S38.14, which can be converted to desired S38.15 (Scheme 38a) using methods reported earlier on. An alternative synthesis of compound S38.14 is also shown in Scheme 38b. Commercially available 2-amino-2-methyl-1-propanol is converted to aziridines S38.16 according to literature methods (*J. Org. Chem.* 1992, 57, 5813; *Syn. Lett.* 1997, 8, 893). Aziridine opening with phosphite give S38.17 (*Tetrahedron Lett.* 1980, 21, 1623). Reprotection of S38.17 affords S38.14.

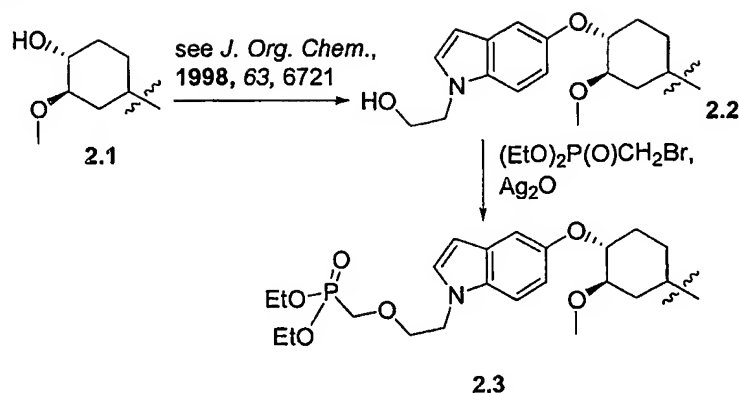
Scheme 38a**Scheme 38b**

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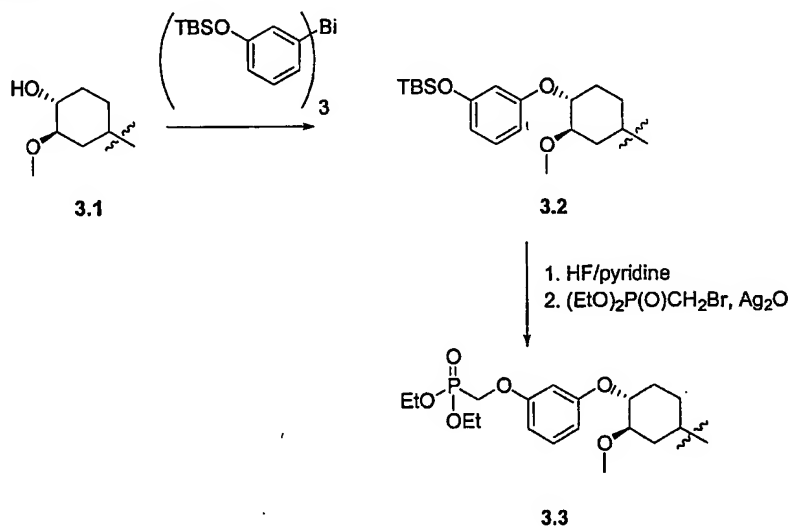
The invention will now be illustrated by the following non-limiting Examples.

Example 1

Rapamycin (compound, **1.1**, wherein the remaining portion of the rapamycin structure is not shown), a synthetic precursor of everolimus, is *O*-arylated as shown above using an appropriate aryl bismuth reagent according to a procedure such as that reported in *Bioorg. Med. Chem. Lett*, **1995**, *5*, 1035. 3-(Dimethyl-*t*-butylsilyloxy)bromobenzene is treated either with magnesium in diethyl ether or with butyllithium in tetrahydrofuran, and the resulting organometallic reagent is reacted with bismuth trichloride to generate the triarylbismuthine. After treating with 1 – 1.2 equivalents of peracetic acid, the bismuth(V) reagent is mixed with rapamycin and copper(II) acetate. The reaction is allowed to proceed for a day at room temperature or, if necessary, at reflux, affording the desired 3-(dimethyl-*t*-butylsilyloxy)phenyl ether, **1.2**. After removal of the dimethyl-*t*-butylsilyl protecting group, *O*-alkylation is achieved with diethyl (bromomethyl)phosphonate in the presence of silver oxide, affording the desired everolimus analog containing the diethylphosphonate, **1.3**. Silver ion-assisted reactions have been used to mediate *O*-alkylations on an immunosuppressive macrolide structurally similar to rapamycin: see *J. Med. Chem.*, **1998**, *41*, 1764.

Example 2

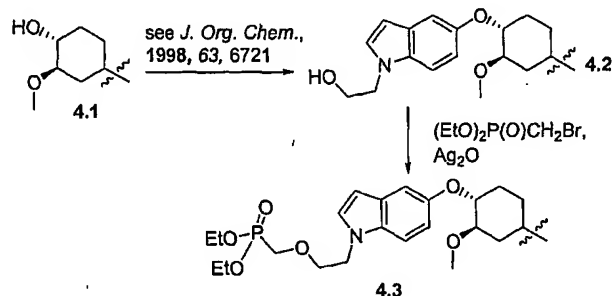
A phosphonate derivative of everolimus indolyl ether is prepared from rapamycin (formula, 2.1, wherein the remaining portion of the rapamycin structure is not shown) in a similar manner to that described in Example 1, with the exception that the key triindolylbismuthine intermediate is obtained from 5-bromoindole following the procedure described in *J. Org. Chem.* 1998, 63, 6721.

Example 3

Tacrolimus (compound, 3.1, wherein the remaining portion of the tacrolimus molecule is not shown) is *O*-arylated as shown above using an appropriate aryl bismuth reagent according to a procedure such as that reported in *Bioorg. Med. Chem. Lett*, 1995, 5, 1035. 3-(Dimethyl-*t*-butylsilyloxy)-bromobenzene is treated either with magnesium in diethyl ether or with

butyllithium in tetrahydrofuran, and the resulting organometallic reagent is reacted with bismuth trichloride to generate the triarylbismuthine. After treating with 1 – 1.2 equivalents of peracetic acid, the bismuth(V) reagent is mixed with tacrolimus, **3.1**, and copper(II) acetate. The reaction is allowed to proceed for a day at room temperature or, if necessary, at reflux, affording the desired 3-(dimethyl-t-butylsilyloxy)phenyl ether. After removal of the dimethyl-t-butylsilyl protecting group with HF, *O*-alkylation is achieved with diethyl (bromomethyl)phosphonate in the presence of silver oxide, affording the desired tacrolimus analog containing the diethylphosphonate, **3.3**. Silver ion-assisted reactions have been used to mediate *O*-alkylations on an immunosuppressive macrolide structurally similar to tacrolimus. (See *J. Med. Chem.*, 1998, 41, 1764.)

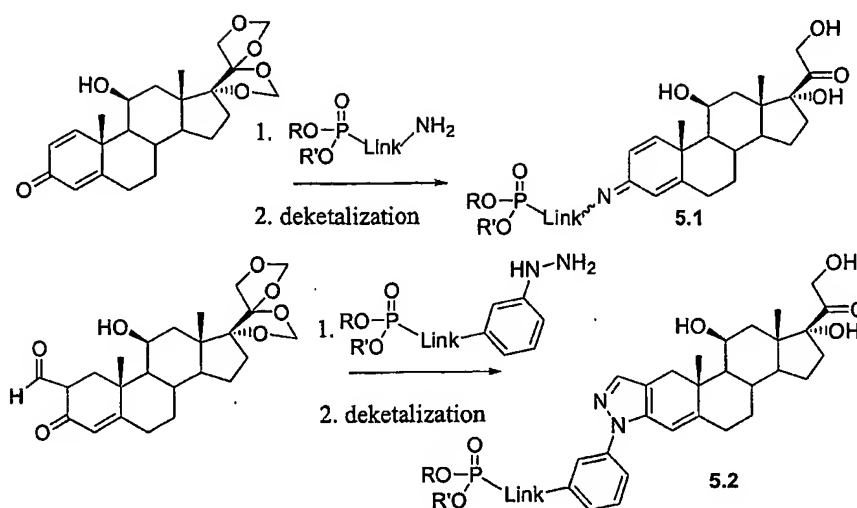
Example 4



A phosphonate derivative of, **4.3**, tacrolimus indolyl ether is prepared from tacrolimus (compound, **4.1**, wherein the remaining portion of the tacrolimus molecule is not shown) in a similar manner to that described in Example 3 with the exception that the key triindolylbismuthine intermediate is obtained from 5-bromoindole following the procedure described in *J. Org. Chem.* 1998, 63, 6721.

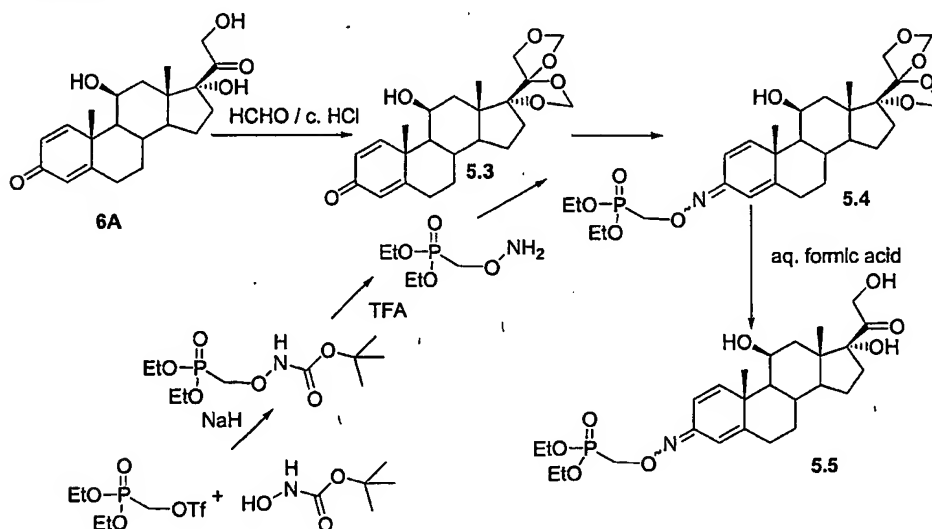
Example 5

Representative compounds of the invention can be made by procedures such as those described by Boer, et al, *J. Mass Spectrom.* 1995, 30, 497-504 and Hoyte, et al, *J. Med. Chem.* 2002, 45, 5397-5405; they can also be made according to the following general routes.



Compounds of formulae, 5.1 and 5.2, wherein "Link" have any of the values defined herein for a linking group or a linker, are representative compounds of the invention.

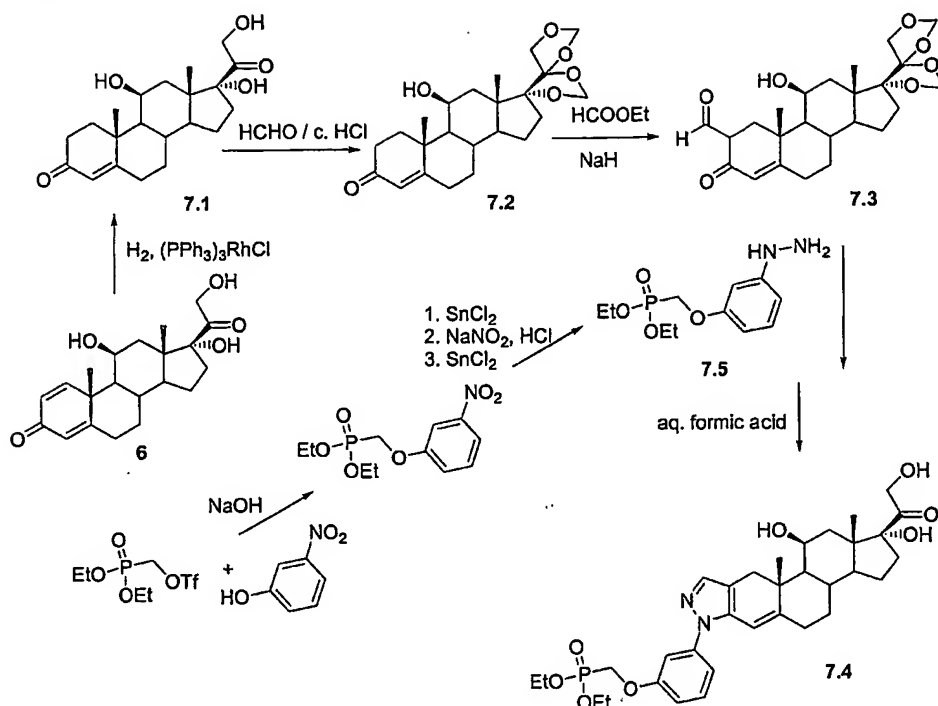
Example 6



Prednisolone, 6A, is treated in a solvent, such as chloroform, with formaldehyde in the presence of an acid, such as concentrated hydrochloric acid. After stirring for several hours (preferably 7 to 10 hours) at room temperature, the layers are separated and the organic layer is concentrated to afford the bis-(methylenedioxy) intermediate, 5.3 (Hirschmann, R. et al, *J. Am. Chem. Soc.*

1964, 86, 1520-1527). This intermediate is treated with diethyl (aminooxymethyl)phosphonate in a solvent such as pyridine to afford the oxime, 5.4. The oxime is treated with aqueous acid to remove the bis-(methylenedioxy) protecting group. For example, the oxime is treated with 60% aqueous formic acid and heated at 90 °C for 10 min., cooled and concentrated using portions of ethanol to assist in removing formic acid. Chromatographic purification and/or crystallization of the residue yield the phosphonate oxime analog, 5.5, of prednisolone.

A key precursor of this synthesis, diethyl (aminooxymethyl)phosphonate, can be obtained from diethyl (trifluoromethyl-sulfonyloxymethyl)phosphonate and N-(t-butoxycarbonyl)-hydroxylamine. Accordingly, N-(t-butoxycarbonyl)-hydroxylamine is dissolved in a solvent such as THF and treated with sodium hydride. When bubbling ceases, diethyl (trifluoromethylsulfonyloxymethyl)phosphonate (prepared according to *Tetrahedron Lett.*, 1986, 27, 1477) is added. After quenching the reaction with aqueous ammonium chloride and extracting the product with an organic solvent such as ethyl acetate, the N-Boc protected diethyl (aminooxymethyl)phosphonate is isolated by chromatography. The N-Boc protecting group is then removed by treatment of trifluoroacetic acid, affording the desired diethyl (aminooxymethyl)phosphonate.

Example 7

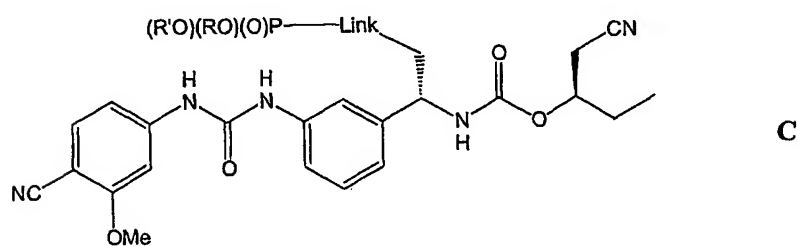
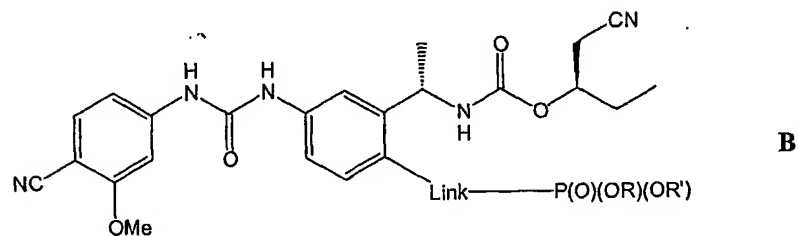
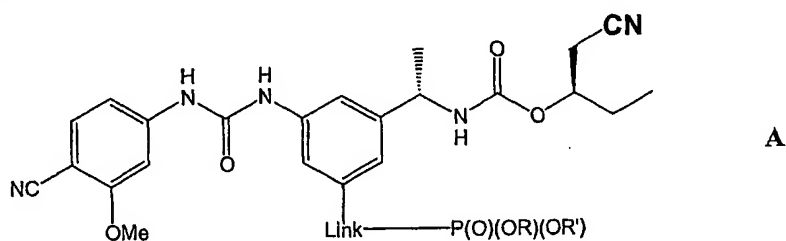
Prednisolone, 6A, is reduced to 1,2-dihydroprednisolone, 7.1, using a rhodium catalysis such as tris(triphenylphosphine)rhodium(I) chloride under hydrogen according to a procedure such as that reported by Procopiu, P. et al, *J. Med. Chem.*, 2001, 44, 602-612. The dihydroxy ketone group on the D ring of the steroid is then protected using the method described in Example 6, before formylation at the C-2 position. For example, the bis-(methylenedioxy) intermediate, 7.2, is treated with freshly distilled ethyl formate and sodium hydride in a solvent such as toluene. The reaction is quenched with aqueous solution of a weak base such as potassium dihydrogen phosphate. The crude product is purified by a general method such as crystallization, affording the 2-formyl intermediate, 7.3. This 2-formyl compound is condensed with a phosphonate-substituted phenylhydrazine to yield, after removal of the bis-(methylenedioxy) protecting group, the desired phosphonate pyrazole analog, 7.4, of prednisolone.

A key precursor, 3-[(diethylphosphono)-methoxy]phenylhydrazine, 7.5, can be made starting from diethyl (trifluoromethylsulfonyloxymethyl)-phosphonate and 3-nitrophenol. 3-Nitrophenol is treated with a base such as

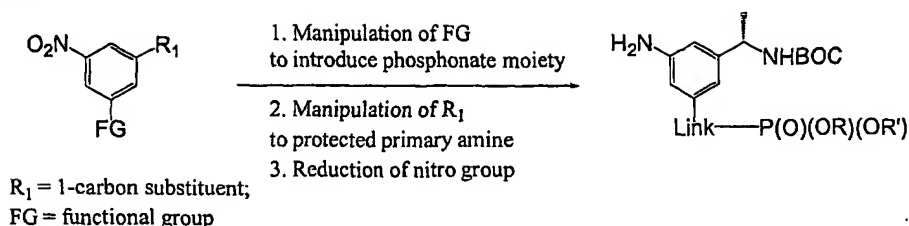
sodium hydroxide and then *O*-alkylated with diethyl (trifluoromethylsulfonyl-oxymethyl)phosphonate. The nitro group is reduced with tin(II) chloride and subsequently converted to the aryl hydrazine by diazotization and reduction with sodium sulfite (*Chem. Ber.*, 1960, 93, 540) or tin(II) chloride (*J. Med. Chem.*, 2001, 44, 4031).

Examples 8-13

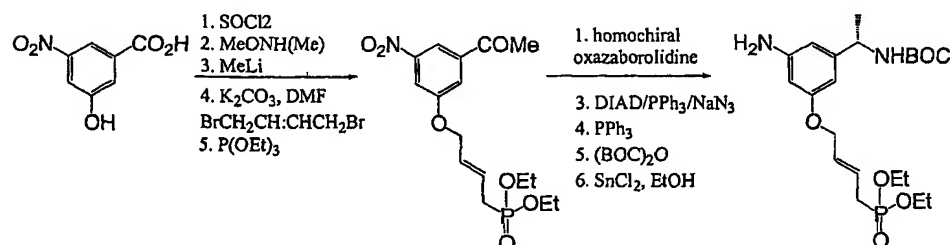
Synthetic methodologies and intermediate compounds that can be used to prepare VX-148 analogs of formulae A, B, or C are described in Examples 8-13. The below compounds are representative examples of compounds of Formulae 6, 7, and 8.



Link includes 0-8 atoms; 2 - 6 is preferred.

Example 8

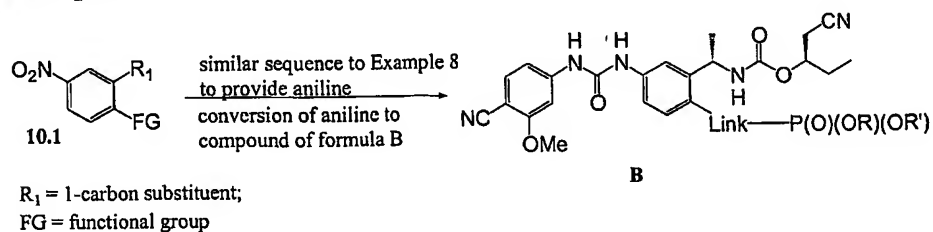
A general scheme that is useful for converting a 3,5-difunctionalized nitrobenzene derivative to an aniline that can be used to prepare a VX-148 analog of the invention is illustrated above.

Example 9

3-Hydroxy-5-nitro-benzoic acid is heated briefly in thionyl chloride to generate the acid chloride. The acid chloride is condensed with O,N-dimethyl-hydroxylamine in the presence of a base such as triethylamine to produce the Weinreb amide which, upon reaction with methyl lithium, provides the acetophenone derivative. The acetophenone derivative is treated with a base such as potassium carbonate in a dipolar aprotic solvent such as dimethyl-formamide, in the presence of an excess of *E*-1,4-dibromobutene. The monobromide is isolated by chromatography and then subjected to treatment with triethylphosphite in a solvent such as toluene (or other Arbuzov reaction conditions: see Engel, R., "Synthesis of Carbon-phosphorus Bonds," CRC press, 1988) to generate the desired phosphonate diethyl ester. Thereafter, the carbonyl of the acetophenone is reduced enantioselectively using an appropriate homochiral oxazaborolidine such as that described by Corey (*J. Am. Chem. Soc.*, 1987, 109, 5551), and the resulting alcohol is displaced by azide using a method such as that described by Mitsunobu (*Bull. Chem. Soc. Japan.*, 1971, 44, 3427). The azide is reduced to the amine under Staudinger conditions (*Helv. Chim. Act.*, 1919, 2, 635) and protected as the *t*-butyl carbonate. Finally, the desired aniline

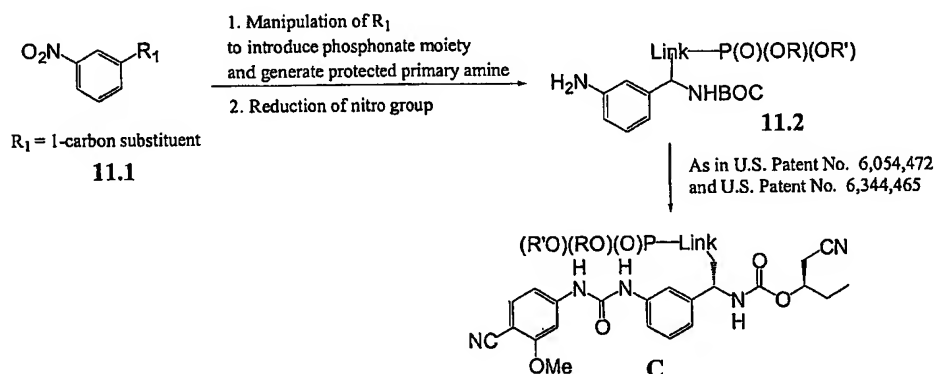
intermediate is generated by tin (II)-mediated reduction of the nitrobenzene. The aniline is converted to a compound of Formula A (also Formula 6) using coupling reactions similar to those described in U.S. Patent No. 6,054,472 and U.S. Patent No. 6,344,465.

Example 10

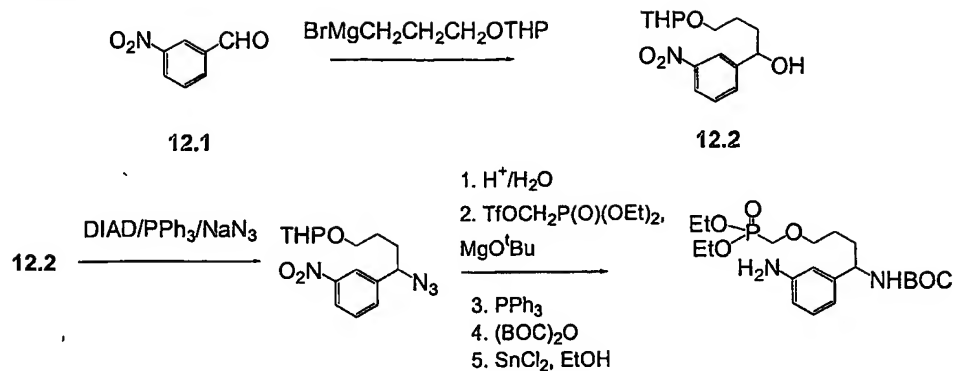


A general scheme that is useful for converting a 3,4-disubstituted nitrobenzene derivative, 10.1, to an aniline, which can be converted to a compound of Formula B (also Formula 7) using coupling reactions similar to those described in U.S. Patent No. 6,054,472 and U.S. Patent No. 6,344,465, is illustrated above.

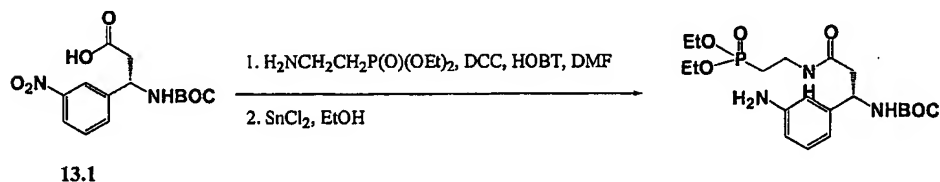
Example 11



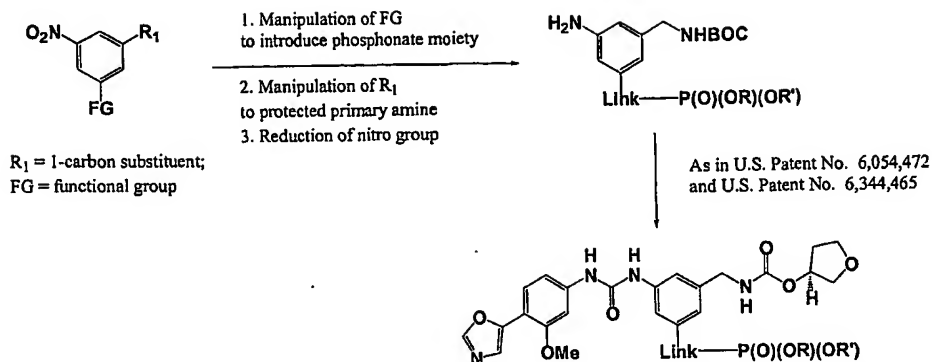
Manipulation of a 3-substituted nitrobenzene, 11.1, provides aniline, 11.2, which can be converted to a compound of formula C (also Formula 8) using coupling reactions similar to those described in U.S. Patent No. 6,054,472 and U.S. Patent No. 6,344,465.

Example 12

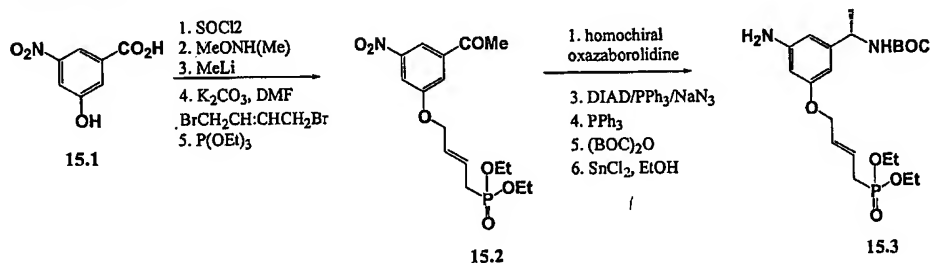
3-Nitrobenzaldehyde, 12.1, reacts with a Grignard reagent to introduce a tether bearing a protected alcohol and simultaneously to generate a benzylic alcohol, as shown. The alcohol, 12.2, is displaced by an azide in a manner similar to that described for Example 9. After deprotection, the liberated alcohol is alkylated with diethyl phosphonomethyltriflate (prepared according to *Tetrahedron Lett.*, 1986, 27, 1477) using a base such as magnesium tert-butoxide in a solvent such as tetrahydrofuran. Subsequent transformations of the azide and nitro groups proceed in a fashion similar to that described in Example 9. See Batt et al, *Bioorg. Med. Chem. Lett.*, 1995, 5, 1549.

Example 13

3-tert-Butoxycarbonylamino-3-(3-nitro-phenyl)-propionic acid, 13.1, (commercially available) is coupled with 2-aminoethylphosphonic acid diethyl ester (commercially available) using standard reagents for the formation of a secondary amide such as dicyclohexylcarbodiimide (DCC) and hydroxy-benzotriazole (HOBT), in a solvent such as dimethylformamide. Subsequent reduction of the nitro group proceeds in a fashion similar to that described in Example 9.

Example 14

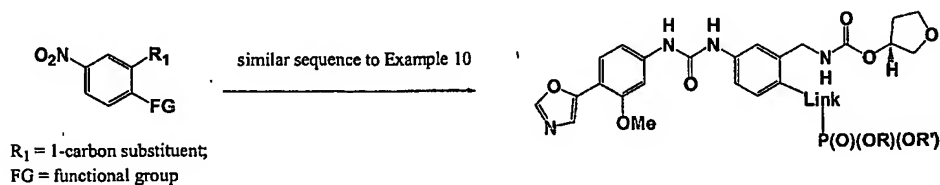
The above scheme illustrates a general route that can be used to prepare compounds of Formula 9.

Example 15

3-Hydroxy-5-nitro-benzoic acid, **15.1**, is heated briefly in thionyl chloride to generate the acid chloride. This is then condensed with *O,N*-dimethyl-hydroxylamine in the presence of a base such as triethylamine to produce the Weinreb amide which, upon reaction with methyl lithium, gives the acetophenone derivative. This is then treated with a base such as potassium carbonate in a dipolar aprotic solvent such as dimethyl-formamide, in the presence of an excess of *E*-1,4-dibromobutene. The monobromide is isolated by chromatography and then subjected to treatment with triethylphosphite in a solvent such as toluene (or other Arbuzov reaction conditions: see Engel, R., "Synthesis of Carbon-phosphorus Bonds," CRC press, 1988) to generate the desired phosphonate diethyl ester, **15.2**. Thereafter, the carbonyl of the acetophenone is reduced enantioselectively using an appropriate homochiral oxazaborolidine such as that described by Corey (*J. Am. Chem. Soc.*, 1987, 109, 5551), and the resulting alcohol is displaced by azide using a method such as that described by Mitsunobu (*Bull. Chem. Soc. Japan.*, 1971, 44, 3427). The azide is

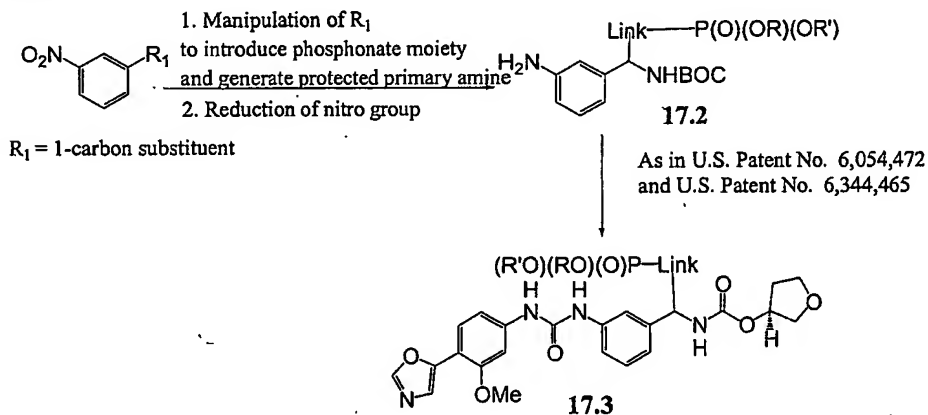
reduced to the amine under Staudinger conditions (*Helv. Chim. Act.*, 1919, 2, 635) and protected as the t-butyl carbonate. Finally, the desired aniline intermediate, 15.3, is generated by tin (II)-mediated reduction of the nitrobenzene.

Example 16



Reagents suitable for use in the synthesis of representative compounds of Formula 10 may be made by routes analogous to that shown in Example 10, starting from 2-hydroxy-5-nitro-benzoic acid.

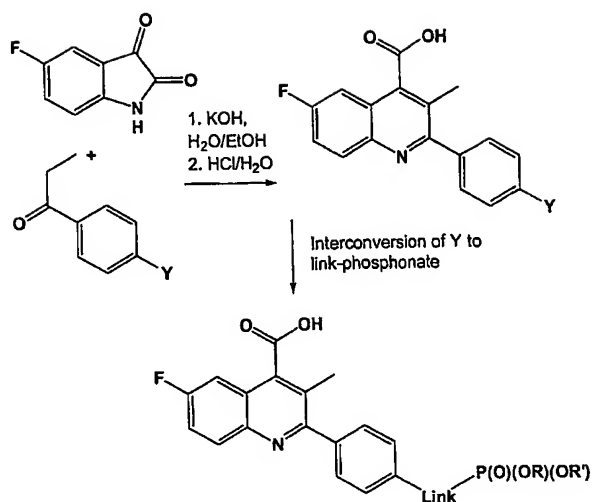
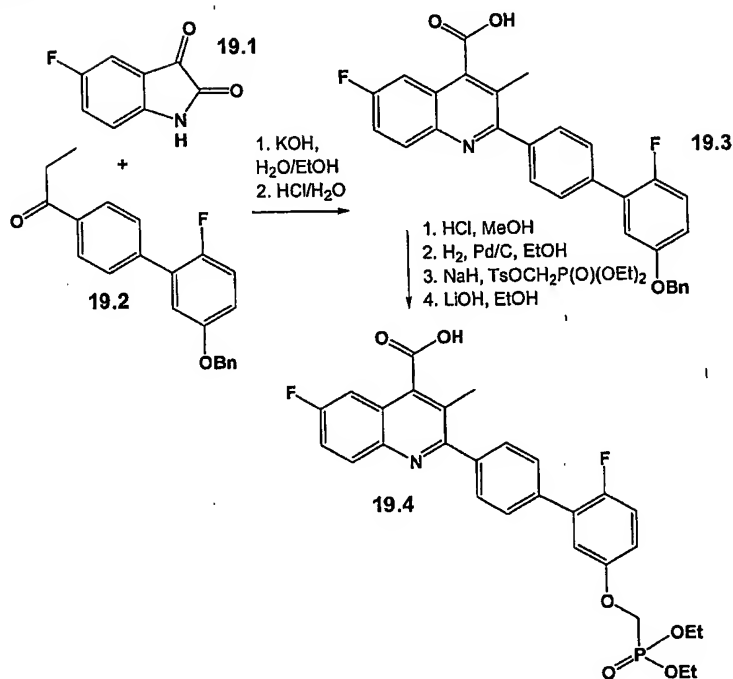
Example 17



Representative compounds of Formula 11 can be prepared as illustrated in Examples 11-13, above. The preparation of anilines of formula, 17.2, is illustrated in Examples 11-13 above. Anilines of formula, 17.2, can be converted to compounds of Formula 11 using procedures similar to those described in U.S. Patent No. 6,054,472 and U.S. Patent No. 6,344,465.

Example 18

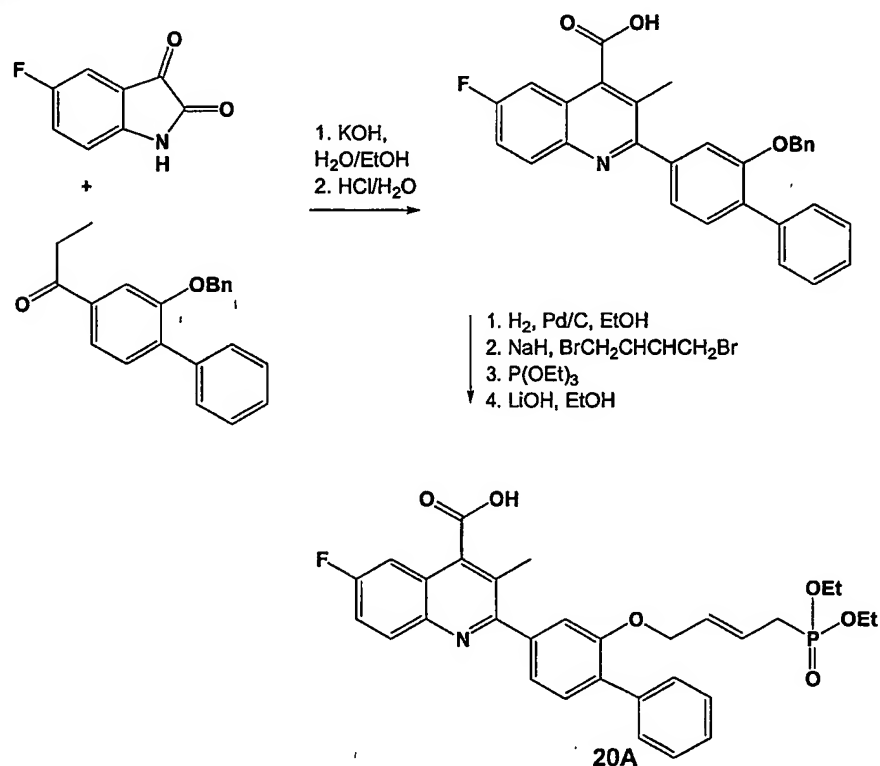
The following is a general route that can be used to prepare compounds of Formula 15.

**Example 19**

The initial Pfitzinger condensation of compound 19.1 and compound 19.2 is achieved in a single step using potassium hydroxide with acidic work-up, as shown. Alternatively, the initial aldol condensation may be performed using diethylamine in ethanol, and the quinoline ring may be formed in a second step mediated by an acid such as hydrochloric acid in a solvent such as 1,4-dioxane. Following removal of the benzyl protecting group via hydrogenation, the phenol

can be treated in a solvent such as tetrahydrofuran or dimethylformamide with a base such as sodium hydride. When bubbling ceases, diethyl phosphonomethyltriflate (prepared according to *Tetrahedron Lett.*, 1986, 27, 1477) is added, yielding the desired phosphonate diester. The carboxylate is deprotected by treatment with lithium hydroxide in ethanol to provide compound 19.4 (which is a compound of Formula 12).

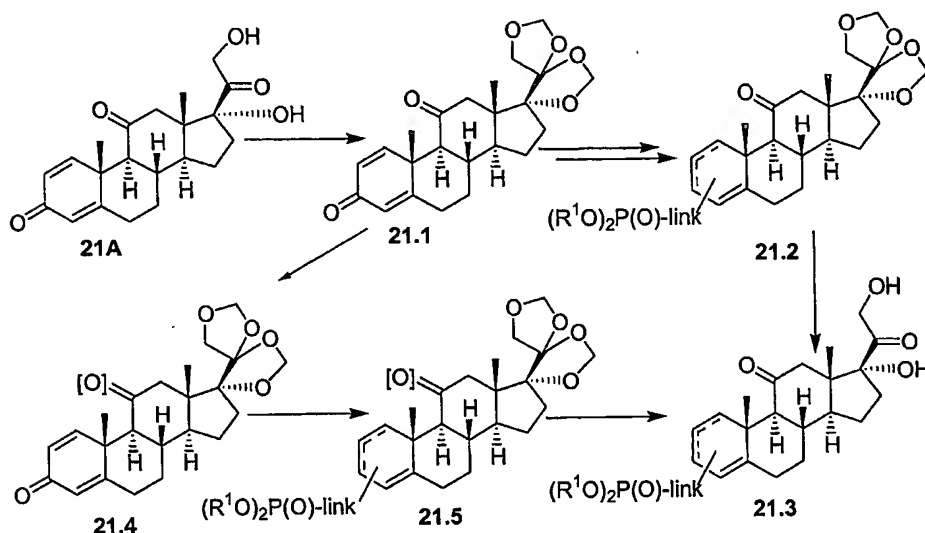
Example 20



The synthesis is similar to that depicted in Example 19 except that, following deprotonation of the phenol, *E*-1,4-dibromobutene is added in excess. After quenching the reaction with aqueous ammonium chloride and extracting the product with an organic solvent such as ethyl acetate, the mono-alkylated product is isolated by chromatography. The resulting bromide is heated with triethylphosphite in a solvent such as toluene to generate the diethyl ester of the desired phosphonic acid, and the carboxylic acid is deprotected as before to provide compound 20A (which is a compound of Formula 12).

Example 21

The structures of Prednisone, **21A** (U.S. Patent No. 2,897,464), and representative phosphonate esters are shown below, in which the substituent R^1 is H, alkyl, alkenyl, aryl or aralkyl. The phosphonate compounds incorporate a phosphonate moiety $(R^1O)_2P(O)$ connected to the nucleus by means of a variable linking group, designated as "link."



For example, the above scheme depicts a protection-deprotection sequence in which the steroid side-chain is protected as a bis-methylenedioxy (BMD) moiety. In this sequence, Prednisone, **21A**, is reacted with paraformaldehyde and an acid catalyst such as hydrochloric acid, as described in "Protective Groups in Organic Synthesis," by T.W. Greene and P.G.M. Wuts, Wiley, Second Edition 1990, p. 223, to yield the BMD derivative, **21.1**. The phosphonate moiety is then introduced, using the procedures described below, to produce the phosphonate ester, **21.2**. The BMD moiety is then hydrolyzed, for example by treatment with 50% aqueous acetic acid, as described in "Protective Groups in Organic Synthesis," by T.W. Greene and P.G.M. Wuts, Wiley, Second Edition 1990, p. 223, to afford the triol, **21.3** (which is a compound of Formula 23).

Optionally, depending on the nature of the reactions employed, the 11-ketone group in the BMD compound, **21.1**, is protected before introduction of the phosphonate group. The ketone is protected, for example, as the cyclic

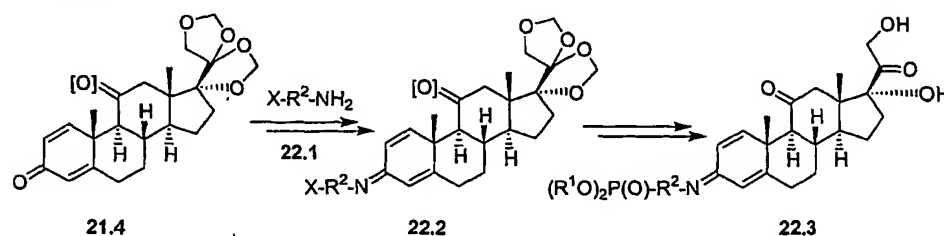
ethylene ketal, by reaction in toluene solution at reflux temperature with ethylene glycol and an acid catalyst, as described in *J. Am. Chem. Soc.*, **1955**, 77, 1904. Deprotection is effected by reaction with pyridinium tosylate in aqueous acetone, as described in *J. Chem. Soc., Chem. Comm.*, 1351, 1987.

Alternatively, the 11-ketone is protected by conversion to the N, N-dimethylhydrazone. The dimethyl hydrazone is prepared by the reaction of the ketone, **21.1**, with N,N-dimethylhydrazine in ethanol-acetic acid, as described in *Org. Syn.*, **1970**, 50, 102. The group is removed by treatment with sodium acetate and acetic acid in aqueous tetrahydrofuran, as described in *J. Am. Chem. Soc.*, **1979**, 101, 5841.

Alternatively, the 11-ketone is protected as the diethylamine adduct. In this procedure, the substrate, **21.1**, is reacted with titanium tetrakis-(diethylamide), as described in *J. Chem. Soc., Chem. Comm.*, 406, 1983, to afford the adduct. The ketone is deprotected by reaction with water in an aqueous organic solvent.

The 11-protected BMD compound, **21.4**, is then converted, using the procedures described below, into the phosphonate, **21.5**. Deprotection then yields the 11-keto diol, **21.3**.

Example 22

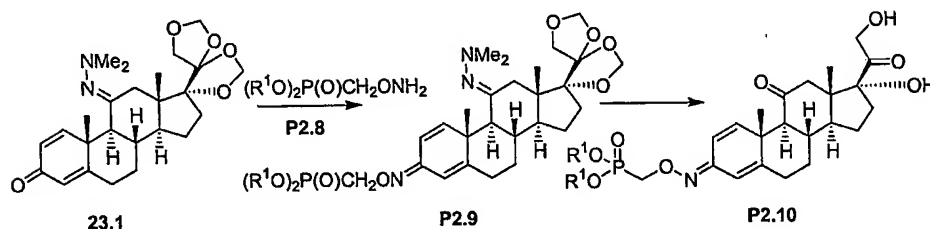


The preparation of phosphonates, **22.3**, in which the phosphonate is attached by means of an imino or iminoxy group and a variable carbon chain are depicted above. In this procedure, the doubly-protected derivative, **21.4**, is reacted with an amine or hydroxylamine, **22.1**, in which R^2 is an alkyl, alkenyl, cycloalkyl or cycloalkenyl group, optionally incorporating a heteroatom O, S or N, or a functional group such as an amide, ester, oxime, sulfoxide or sulfone etc, or an optionally substituted aryl, heteroaryl or aralkyl group, optionally

incorporating a heteroatom O, S or N, and X is either a phosphonate group or a group which is subsequently converted into a phosphonate-containing substituent.

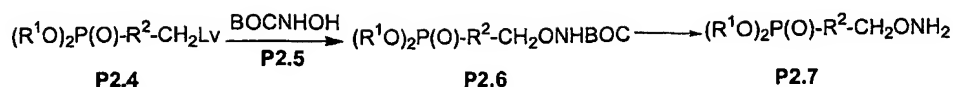
For example, X is dialkylphosphono, bromo, hydroxy, amino, carboxy and the like. The reaction is conducted between equimolar amounts of the reactants in an aprotic solvent such as pyridine or xylene, or in an alcoholic solvent such as ethanol, optionally in the presence of an acid catalyst, to give the imine or oxime. The preparation of oximes of steroidal 3-ketones is described in *Anal. Bioch.*, 1978, 86, 133 and in *J. Mass. Spectrom.*, 1995, 30, 497. The protecting groups are then removed to afford the ketodiol, 22.3 (which is a compound of Formula 16).

Example 23



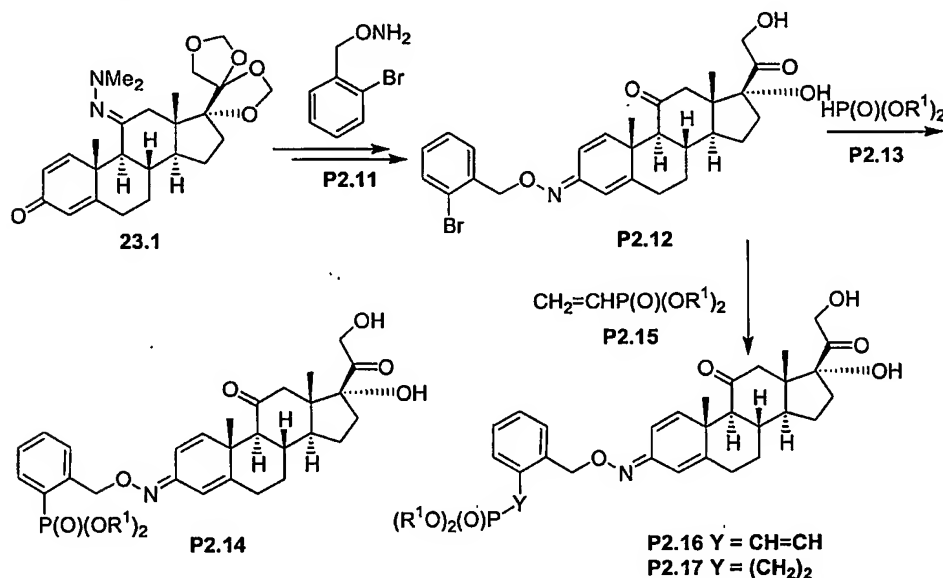
The preparation of phosphonates, **P2.10**, in which the phosphonate is attached by means of an iminoxy group, is illustrated above. The substrate, **23.1**, a compound of Formula 23.1, in which the 11-ketone is protected as the dimethyl hydrazone, is reacted with a dialkyl phosphonomethyl hydroxylamine, **P2.8**, prepared from a dialkyl trifluoromethyl-sulfonyloxymethyl phosphonate (*Tetrahedron Lett.*, 1986, 27, 1477) and BOC-hydroxylamine, to afford the oxime, **P2.9**, which is deprotected by reaction with 50% aqueous acetic acid, to afford the diol, **P2.10**. The oxime forming reaction is typically performed at ambient temperature in ethanol-acetic acid solution between equimolar amounts of the reactants.

The intermediate dialkyl phosphonomethyl hydroxylamine, **P2.8**, (compound, 2.7, wherein R^2 is a bond) can be prepared as follows.



A phosphonate, **P2.4**, in which Lv is a leaving group such as bromo or trifluoromethylsulfonyloxy, is reacted with BOC-hydroxylamine, **P2.5**, (Aldrich) to produce the ether, **P2.6**. The reaction is typically conducted between equimolar amounts of the reactants in a polar solvent such as dimethylformamide or tetrahydrofuran, in the presence of a base such as potassium hydroxide or dimethylaminopyridine. Deprotection, for example, by treatment with trifluoroacetic acid, then gives the hydroxylamine ether, **P2.7**.

Example 24

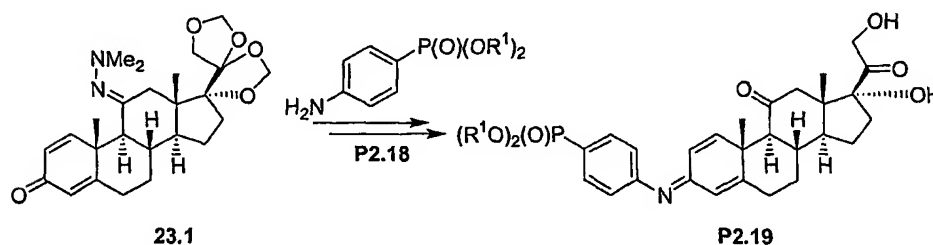


The dienone, **23.1**, is reacted, as described in Example 23, with O-(2-bromobenzyl)hydroxylamine, **P2.11**, prepared from 2-bromobenzyl bromide, to give, after deprotection, the oxime, **P2.12**. The product is then reacted, in the presence of a palladium catalyst, with a dialkyl phosphite, **P2.13**, to afford the phosphonate, **P2.14**. The preparation of arylphosphonates by means of a coupling reaction between aryl bromides and dialkyl phosphites is described in *J. Med. Chem.*, 1992, 35, 1371. The reaction is typically performed in an inert solvent such as toluene, in the presence of a base such as triethylamine and a catalytic amount of tetrakis(triphenylphosphine)palladium(0).

Alternatively, the bromo compound, **P2.12**, is coupled with a dialkyl vinylphosphonate, **P2.15** (Aldrich), to afford the phosphonate, **P2.16**. The coupling of aryl halides with olefins by means of the Heck reaction is described, for example, in "Advanced Organic Chemistry," by F. A. Carey and R. J. Sundberg, Plenum, 2001, p. 503ff and in *Acc. Chem. Res.*, 1979, 12, 146. The aryl bromide and the olefin are coupled in a polar solvent such as dimethylformamide or dioxan, in the presence of a palladium(0) catalyst such as tetrakis(triphenylphosphine)palladium(0) or palladium(II) catalyst such as palladium(II) acetate, and optionally in the presence of a base such as triethylamine or potassium carbonate. Optionally, the styrenoid double bond present in the product, **P2.16**, is reduced, for example by reaction with diimide, to produce the saturated analog, **P2.17**. The reduction of olefinic bonds is described in "Comprehensive Organic Transformations," by R. C. Larock, VCH, 1989, p. 6ff. The transformation is effected by means of catalytic hydrogenation, for example using a palladium on carbon catalyst and hydrogen or a hydrogen donor, or by the use of diimide or diborane.

Using the above procedures, but employing, in place of the benzyloxy reagent, **P2.11**, different bromo-substituted aryl or heteroaryl alkoxy hydroxylamines, and/or different dialkyl alkenyl phosphonates, the products analogous to the compounds, **P2.14**, **P2.16** and **P2.17** are obtained.

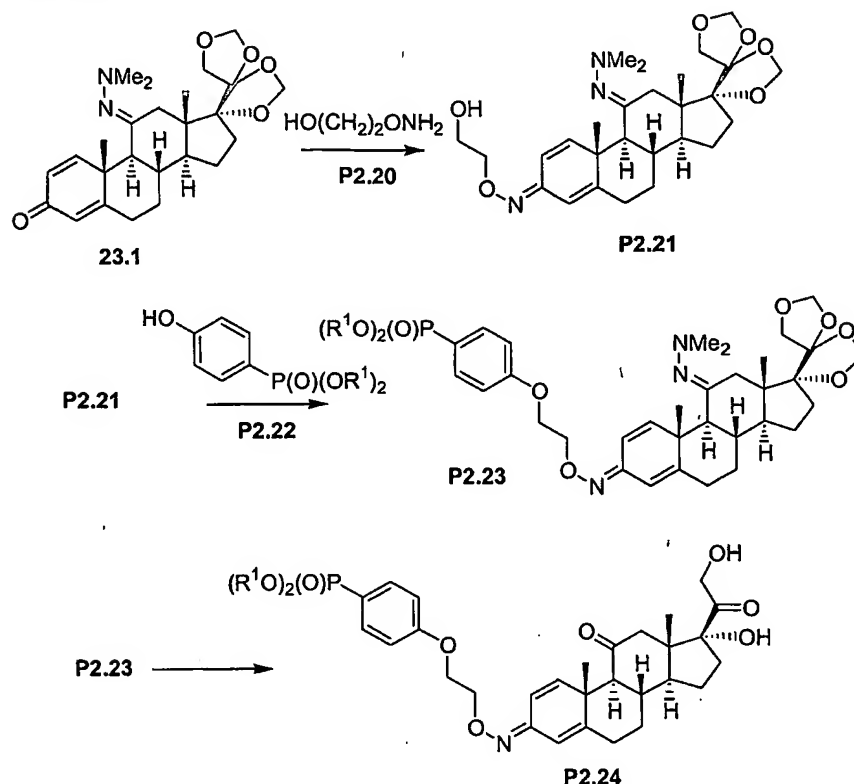
Example 25



The preparation of phosphonates of Formula 16 wherein the phosphonate is attached by means of an imino group is illustrated above. The substrate, **23.1**, is reacted with a dialkyl 4-aminophenyl phosphonate, **P2.18** (Epsilon), to give, after deprotection, the imine product, **P2.19**. The reaction is typically conducted in a hydrocarbon solvent such as toluene or xylene, at reflux temperature, in the

presence of a basic catalyst such as sodium methoxide, or an acid catalyst such as p-toluenesulfonic acid, under azeotropic conditions.

Example 26

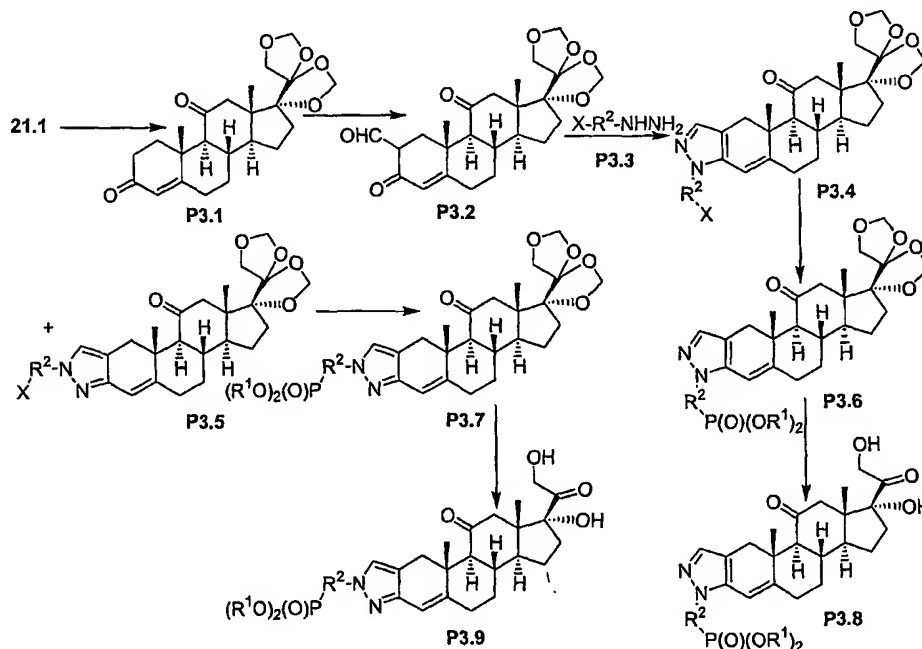


The preparation of representative phosphonates of Formula 16 wherein the phosphonate is attached by means of an oximino group and an ether linkage is illustrated above. In this procedure, the dienone, 23.1, is reacted with O-(2-hydroxyethyl)hydroxylamine, P2.20 (*J. Chem. Soc., Chem. Comm.*, 1986, 903), to yield the oxime, P2.21. The reaction of steroidal 1,4-dien-3-ones with substituted hydroxylamines is described in *J. Steroid Bioch.*, 1976, 7, 795. The reaction is performed between equimolar amounts of the reactants in a polar organic solvent such as pyridine or methanol, optionally in the presence of acetic acid or sodium acetate. The oxime is then reacted in a Mitsunobu reaction with a dialkyl 4-hydroxyphenyl phosphonate, P2.22 (Epsilon), to yield the ether oxime, P2.23. The preparation of aromatic ethers by means of the Mitsunobu reaction is described, for example, in "Comprehensive Organic

Transformations," by R. C. Larock, VCH, 1989, p. 448, and in "Advanced Organic Chemistry," Part B, by F.A. Carey and R. J. Sundberg, Plenum, 2001, p. 153-4 and in *Org. React.*, 1992, 42, 335. The phenol and the alcohol or thiol component are reacted together in an aprotic solvent such as, for example, tetrahydrofuran, in the presence of a dialkyl azodicarboxylate and a triarylphosphine, to afford the ether or thioether products. The procedure is also described in *Org. React.*, 1992, 42, 335-656. The ether product, **P2.23**, is then converted into the ketodiols, **P2.24** (which is a compound of Formula 16).

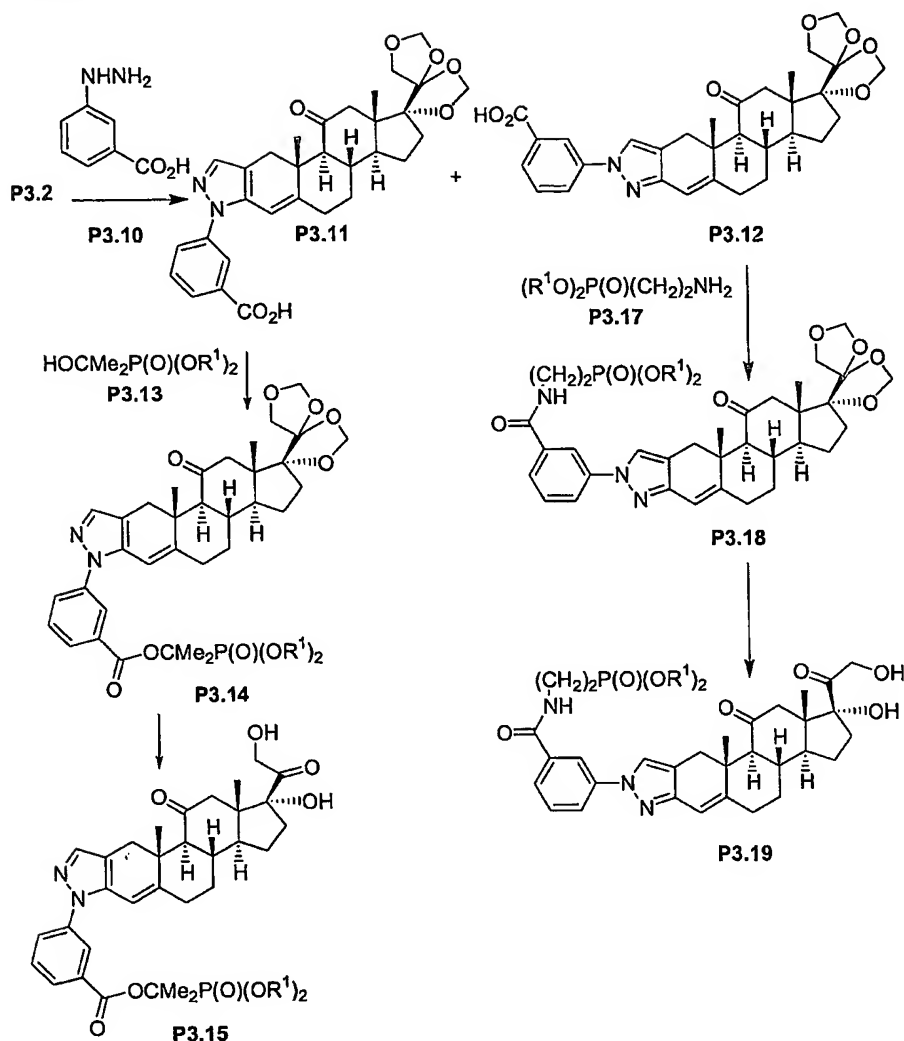
Using the above procedures, but employing, in place of the hydroxylamine, **P2.20**, different hydroxy-substituted hydroxylamines, and/or different hydroxy-substituted aryl phosphonates, the products analogous to **P2.24** are obtained.

Example 27



The preparation of the phosphonate esters of Formulae 17 and 18 in which the phosphonate group is attached to the 1' or 2' position of the pyrazole ring, by means of an aromatic or heteroaromatic group, a heteroatom and a variable carbon chain. In this procedure, the BMD-protected dienone, **21.1**, is reduced to afford the 1,2-dihydro product, **P3.1**. The catalytic hydrogenation reaction is effected by the use of tris(triphenylphosphine)rhodium (I) chloride,

for example as described in *J. Med. Chem.*, **2001**, *44*, 602. The product is then reacted with ethyl formate and a base such as sodium hydride, in an inert solvent such as toluene or dimethylformamide, as described in *J. Am. Chem. Soc.*, **1964**, *86*, 1520, to afford the 2-formyl product, **P3.2**. This compound is then reacted with an alkyl, aralkyl, aryl or heteroaryl hydrazine, **P3.3**, in which the substituent X is either a phosphonate group or a group which is subsequently transformed into a phosphonate-containing substituent. For example, X is dialkylphosphono, bromo, hydroxy, amino, carboxyl and the like. The reaction yields the isomeric 2'- and 1'-aryl pyrazoles, **P3.4** and **P3.5**. The pyrazole-forming reaction is performed between equimolar amounts of the reactants in an acidic solvent such as acetic acid, as described in *J. Am. Chem. Soc.*, **1964**, *86*, 1520. The pyrazoles, **P3.4** and **P3.5**, are then transformed via the BMD-protected intermediates, **P3.6** and **P3.7**, into the phosphonates, **P3.8** and **P3.9**, which are compounds of Formulae **17** and **18** respectively.

Example 28

The preparation of phosphonates in which the phosphonate is attached by means of a phenyl ring and an ester or an amide linkage are illustrated above.

The ketoaldehyde, **P3.2**, is reacted with 3-carboxyphenylhydrazine, **P3.10** (Apin), to give the pyrazoles, **P3.11** and **P3.12**. The 2'-substituted isomer, **P3.11**, is then reacted in dichloromethane solution at ambient temperature with one molar equivalent of a dialkyl 2-hydroxy-2-methylpropyl phosphonate, **P3.13** (French Patent No. 2,462,440), and dicyclohexylcarbodiimide, to yield the ester, **P3.14**. The protecting groups are then removed to yield the diol, **P3.15**, which is a compound of Formula 18.

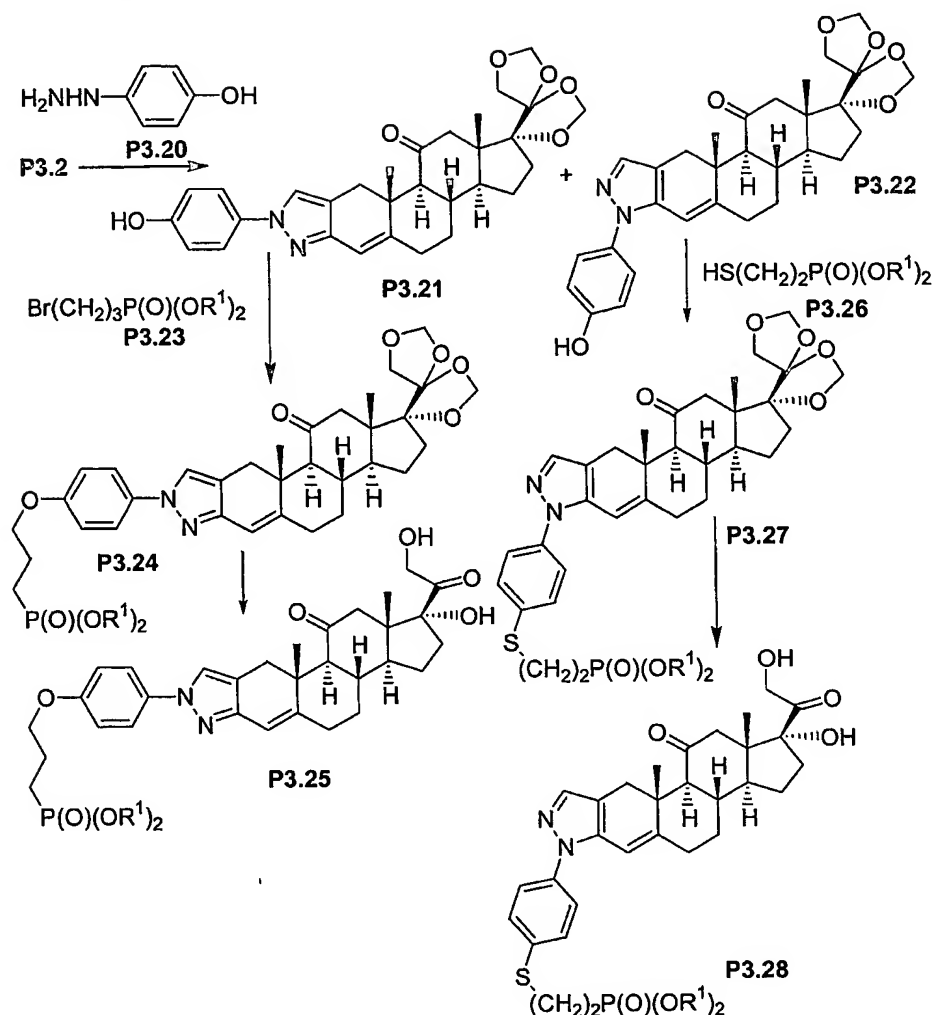
Alternatively, the 1'-substituted pyrazole, **P3.12**, is coupled with a dialkyl 2-aminoethyl phosphonate, **P3.17** (Aurora), to afford the amide, **P3.18**. The

preparation of amides from carboxylic acids and derivatives is described, for example, in "Organic Functional Group Preparations," by S.R.Sandler and W. Karo, Academic Press, 1968, p. 274, and "Comprehensive Organic Transformations," by R.C. Larock, VCH, 1989, p. 972ff. The carboxylic acid is reacted with the amine in the presence of an activating agent, such as, for example, dicyclohexylcarbodiimide or diisopropylcarbodiimide, optionally in the presence of, for example, hydroxybenztriazole, N-hydroxy-succinimide or N-hydroxypyridone, in a non-protic solvent such as, for example, pyridine, DMF or dichloromethane, to afford the amide.

Alternatively, the carboxylic acid may first be converted into an activated derivative such as the acid chloride, anhydride, mixed anhydride, imidazolid and the like, and then reacted with the amine, in the presence of an organic base such as, for example, pyridine, to afford the amide. The conversion of a carboxylic acid into the corresponding acid chloride can be effected by treatment of the carboxylic acid with a reagent such as, for example, thionyl chloride or oxalyl chloride in an inert organic solvent such as dichloromethane, optionally in the presence of a catalytic amount of dimethylformamide. The product, **P3.18**, is then deprotected to give the diol, **P3.19** (which is a compound of Formula 17).

Using the above procedures, but employing different amino or hydroxyl-substituted phosphonates, and/or different carboxy-substituted hydrazines, the products analogous to **P3.15** and **P3.19** are obtained. The functionalization procedures are interchangeable between the pyrazole substrates, **P3.11** and **P3.12**.

Example 29



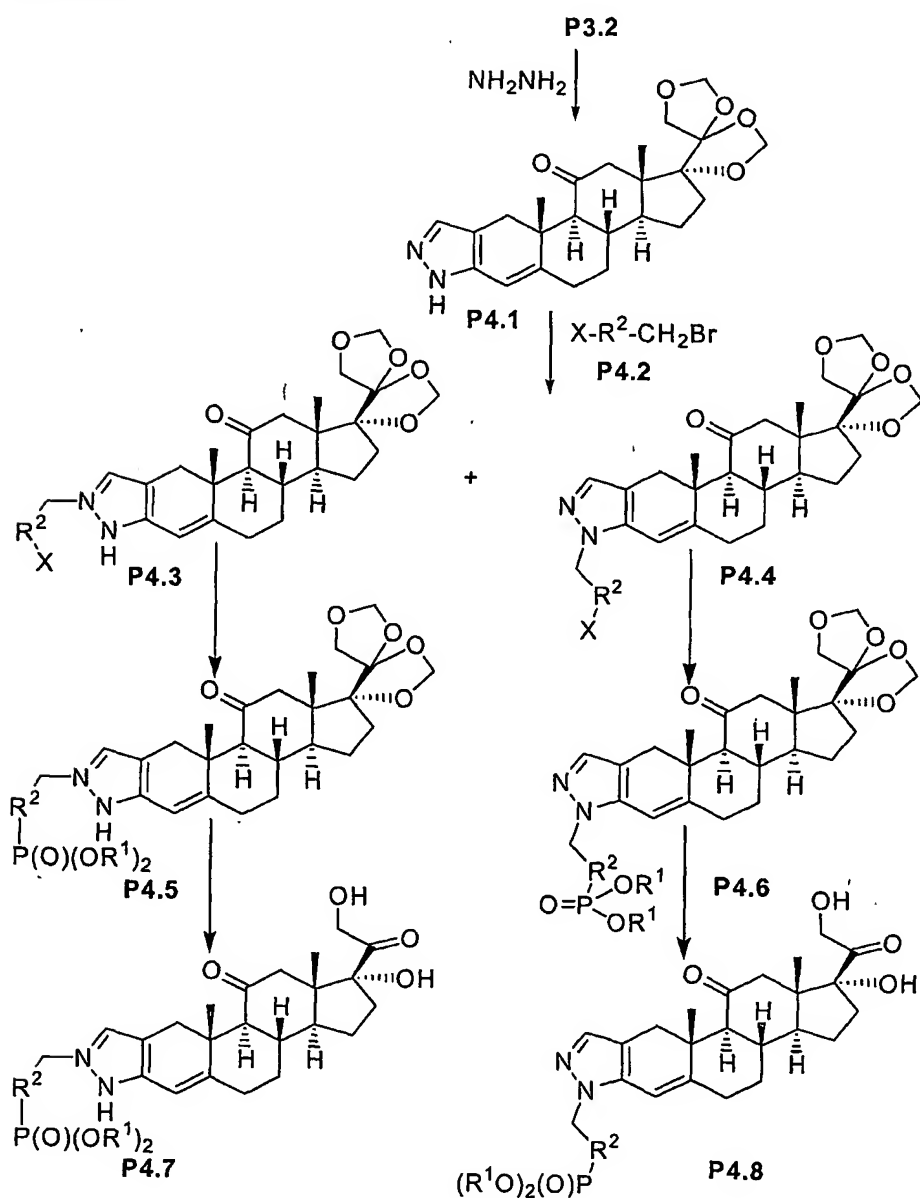
The preparation of the phosphonates of Formulae 17 and 18 wherein the phosphonate group is attached by means of a phenyl group and an alkoxy or alkylthio carbon chain is illustrated above. In this procedure, the ketoaldehyde, **P3.2**, is reacted with 4-hydroxyphenyl hydrazine, **P3.20** (EP 437 105), to produce the pyrazoles, **P3.21** and **P3.22**. The 1'-substituted isomer, **P3.21**, is reacted, in dimethylformamide solution at 70°, with a dialkyl bromopropyl phosphonate, **P3.23** (*J. Amer. Chem. Soc.*, 2000, 122, 1554), and potassium carbonate, to give the phosphonate, **P3.24**. The product is then deprotected to afford the diol, **P3.25**.

Alternatively, the 2'-substituted pyrazole, **P3.22**, is reacted in a Mitsunobu reaction, as described above, with a dialkyl mercaptoethyl

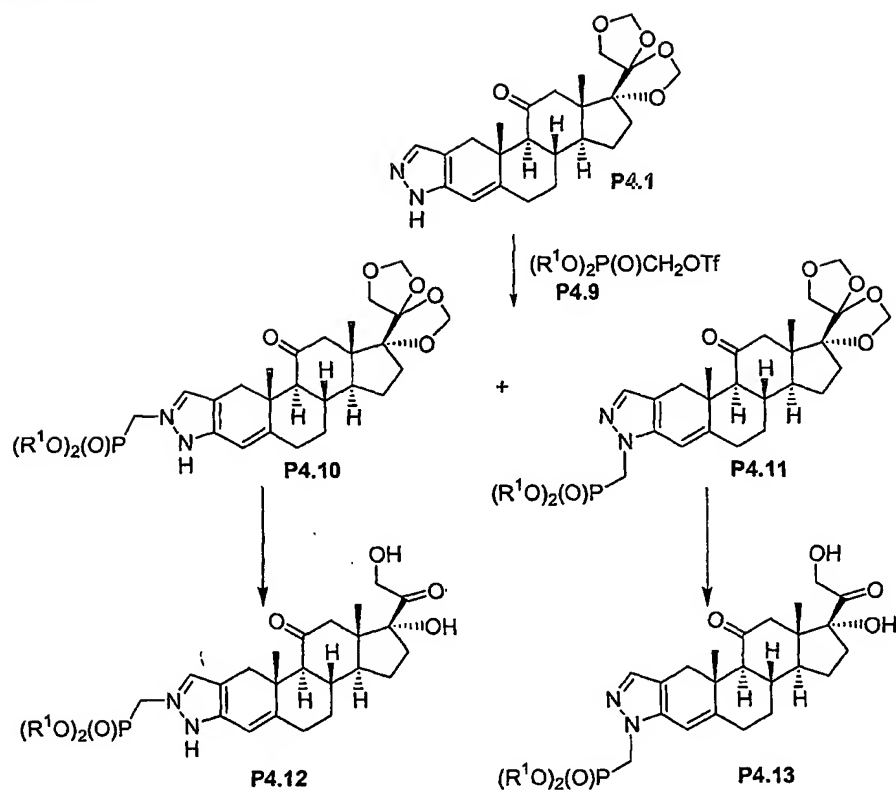
phosphonate, **P3.26** (*Zh. Obshchei. Khim.*, 1973, 43, 2364), to prepare the thioether phosphonate, **P3.27**, which is deprotected to give the diol, **P3.28**.

Using the above procedures, but employing, in place of the hydroxyphenyl reagent, **P3.20**, different hydroxy-substituted aralkyl, aryl or heteroaryl alkoxy hydrazines, and/or different dialkyl bromo or mercapto-substituted phosphonates, the products analogous to the compounds, **P3.25** and **P3.28** are obtained.

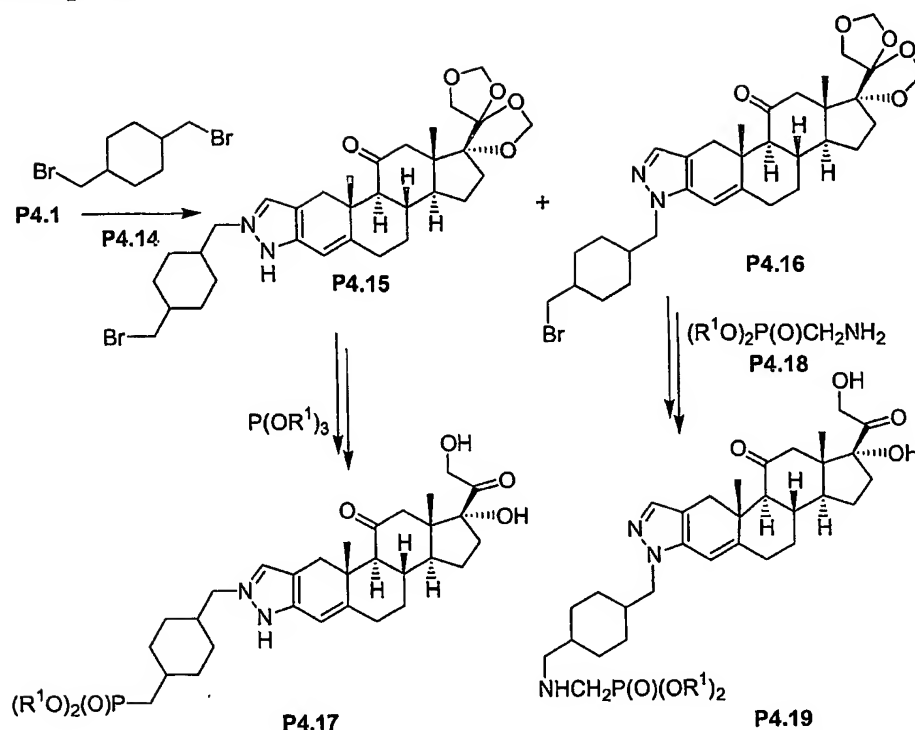
Example 30



The preparation of phosphonate esters of Formulae 17 and 18 wherein the phosphonate group is attached by means of a variable carbon linkage is illustrated above. In this procedure, the ketoaldehyde, **P3.2**, is reacted with hydrazine to afford the pyrazole derivative, **P4.1**. The reaction of steroidal 2-formyl-3-ketones with hydrazine is described in *J. Am. Chem. Soc.*, 1964, 86, 1520. The reaction is performed in acetic acid at ambient temperature. The pyrazole product is then reacted with a bromomethyl compound, **P4.2**, to yield the alkylation products, **P4.3** and **P4.4**. The alkylation of substituted pyrazoles is described, for example, in "Heterocyclic Chemistry," by T. L. Gilchrist, Longman, 1992, p. 309. The reaction is typically performed between equimolar amounts of the substrates in a polar solvent such as dimethylformamide or tetrahydrofuran, in the presence of a base such as dimethylaminopyridine, lithium hexamethyldisilazide and the like. The products, **P4.3** and **P4.4**, are, except in cases where X is dialkylphosphono, converted into the phosphonates, **P4.5** and **P4.6**, using the procedures described herein. Deprotection affords the diols, **P4.7** and **P4.8** (which are compounds of Formulae 18 and 17, respectively).

Example 31

As illustrated above, the pyrazole, P4.1, is reacted with one molar equivalent of a dialkyl trifluoromethanesulfonyloxy phosphonate, P4.9, as described above to give the alkylated pyrazoles, P4.10 and P4.11. Deprotection yields the diols, P4.12 and P4.13 (which are compounds of Formulae 18 and 17 respectively).

Example 32

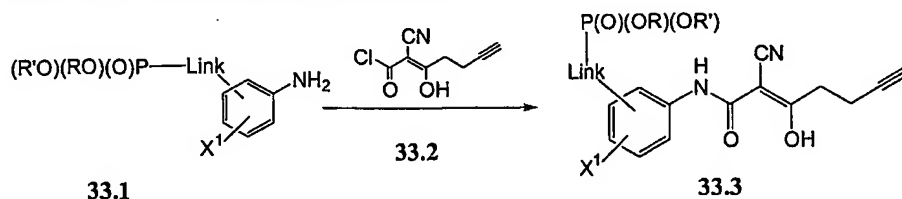
As illustrated above, the pyrazole, **P4.1**, is reacted, as described above, with 1,4-bis(bromomethyl)cyclohexane, **P4.14** (Salor), to give the pyrazoles, **P4.15** and **P4.16**. The product, **P4.15**, is subjected to an Arbuzov reaction, in which the bromomethyl substituent is converted into the dialkyl phosphonomethyl substituent, by reaction with a trialkyl phosphite to prepare, after deprotection of the side chain, the phosphonate, **P4.17**, (which is a compound of Formula 18).

The pyrazole, **P4.16**, is reacted in dimethylformamide with potassium carbonate and a dialkyl aminomethyl phosphonate, **P4.18**, (Interchim) to give after deprotection the amino phosphonate, **P4.19**, which is a compound of Formula 17.

Using the above procedures, but employing, in place of the dibromide, **P4.14**, different dibromides, and/or different amino-substituted phosphonates, the products analogous to **P4.17** and **P4.19** are obtained.

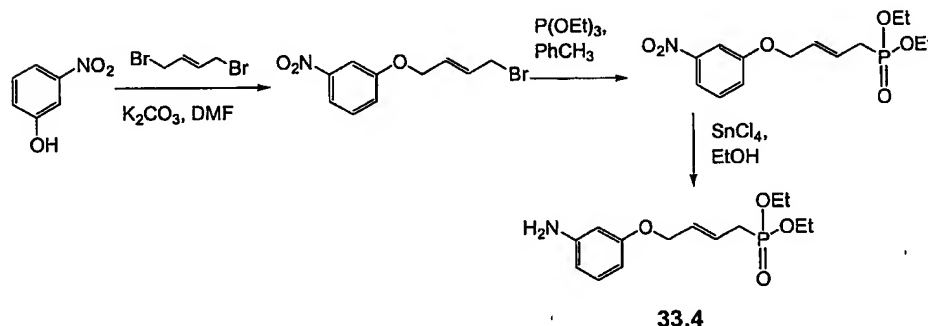
Example 33

Representative compounds of the invention can be prepared as generally described by Westwood et al, *J. Med. Chem.*, 1996, 39, 4608-4621, and according to the following general route.

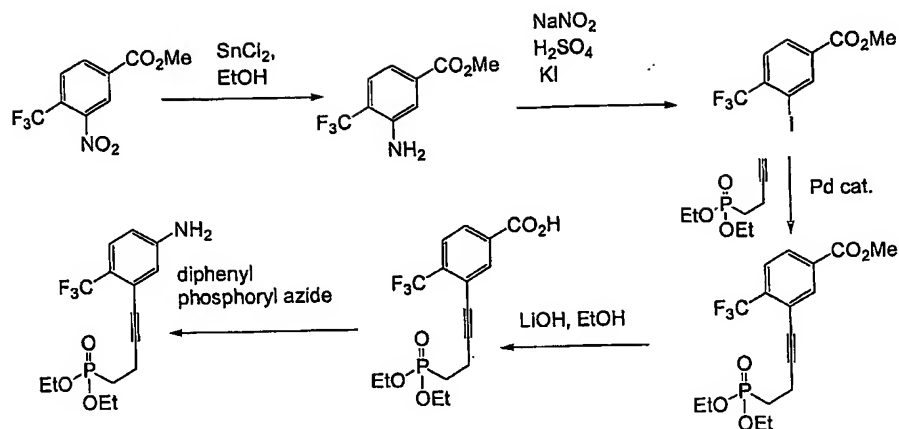


Coupling of a suitable aniline, 33.1, wherein X^1 is hydrogen, halo, trifluoromethyl, (C_1-C_3) alkyl, cyano, or (C_1-C_3) alkoxy, with acid chloride, 33.2, provides compound, 33.3 (which is a representative compound of Formula 19 or 20).

The synthesis of two suitable anilines that can be employed in the above reaction is outlined below.



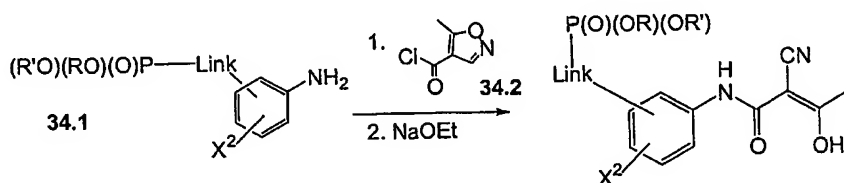
3-Nitrophenol is alkylated with *E*-1,4-dibromobutene and the resulting monobromide is heated with triethylphosphite in a solvent such as toluene (or other Arbuzov reaction conditions) to generate the diethyl ester of the desired phosphonic acid, 33.4. (See Engel, R., "Synthesis of Carbon-phosphorus Bonds," CRC press, 1988). Finally, the desired aniline is generated by tin (II)-mediated reduction of the nitrobenzene.



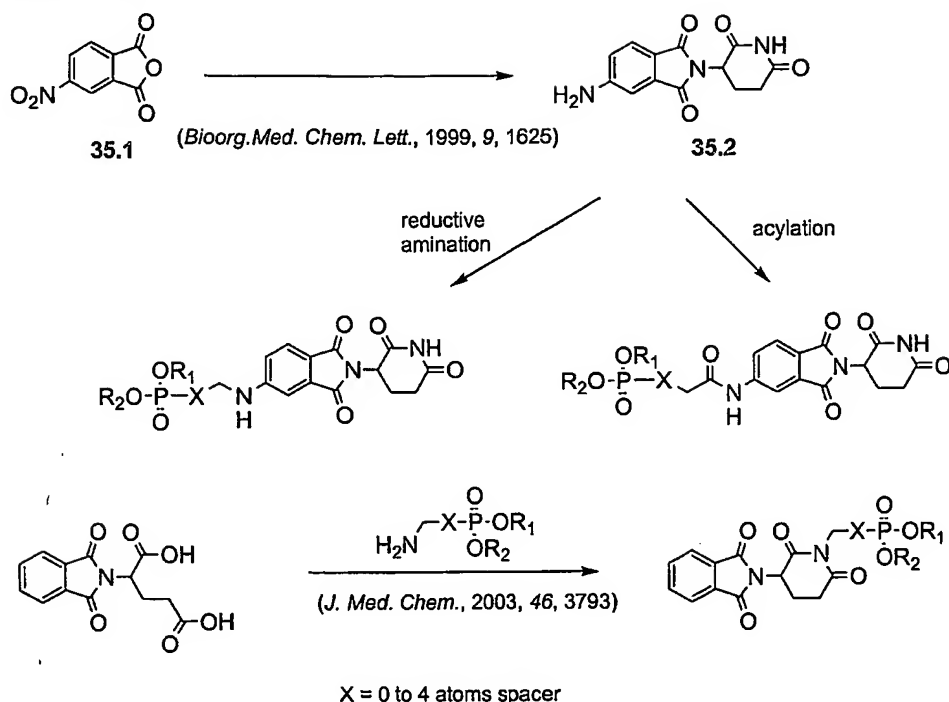
The methyl ester of 3-nitro-4-trifluoromethylbenzoic acid is treated with tin (II) chloride to produce the corresponding aniline. The 3-iodobenzoic acid is generated by diazotization and treatment with potassium iodide. A diethylphosphonate ester is attached via an acetylene linker using palladium catalysis, and after saponification of the benzoate ester, Curtius rearrangement of the acyl azide provides an aniline suitable for incorporation into representative MNA-715 analogs of the invention.

Example 34

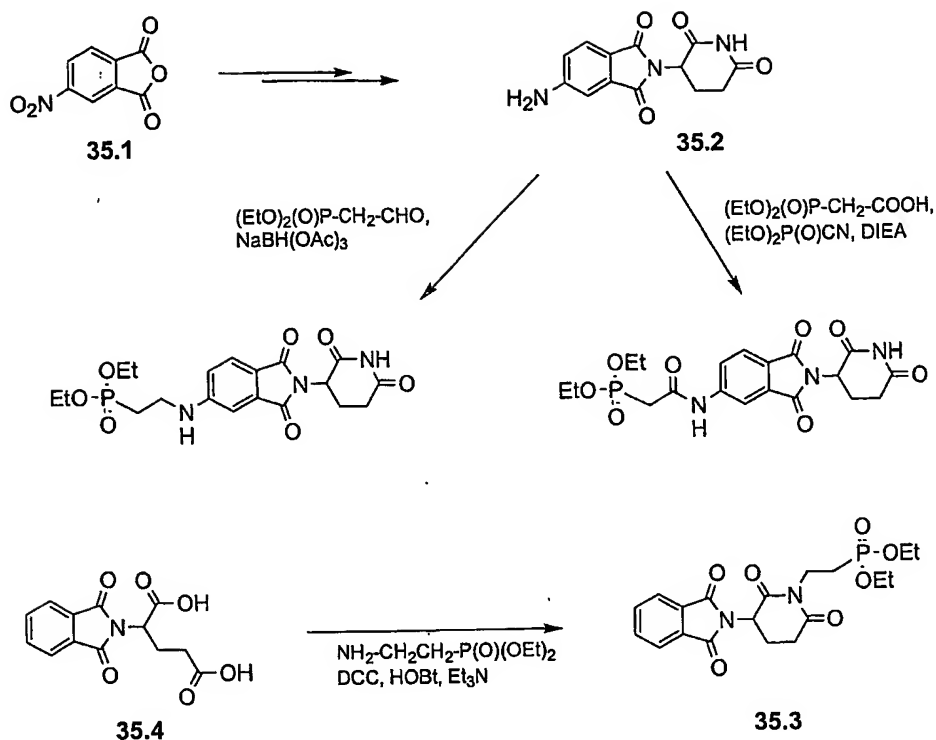
Representative compounds of Formulae 21 and 22 can be prepared as generally described by Westwood et al, *J. Med. Chem.*, 1996, 39, 4608-4621, and according to the following general route.



Coupling of a suitable aniline, 34.1, wherein X^2 is hydrogen, halo, trifluoromethyl, cyano, or methyl with acid chloride, 34.2, provides a representative compound of Formulae 21 and 22. Anilines prepared as described in Example 33 and elsewhere herein can be incorporated into this synthesis.

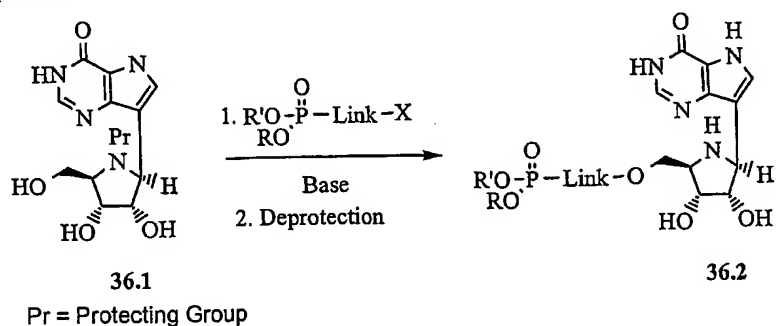
Example 35

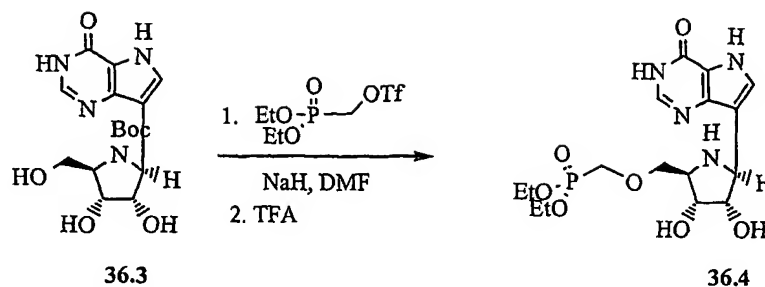
5-Nitro-isobenzofuran-1,3-dione, **35.1**, (commercially available), is converted to 5-amino-2-(2,6-dioxo-piperidin-3-yl)-isoindole-1,3-dione, **35.2**, following the procedures reported in *Bioorg. Med. Chem. Lett.*, 1999, 9, 1625. This amine intermediate is subjected to a reductive amination with diethylphosphonoacetaldehyde (obtained from ozonolysis of diethyl allylphosphonate) in the presence of a reducing agent such as sodium triacetoxyborohydride to generate the desired amine linker analog (*J. Org. Chem.*, 1996, 61, 3849). Alternatively, the amine is acylated with an activated diethylphosphonoacetic acid to provide the desired amide linker compound, according to a procedure such as those reported in *J. Med. Chem.*, 1982, 25, 960 and *J. Med. Chem.*, 1984, 27, 600. The activated diethylphosphonoacetic acid can be obtained by treatment in a solvent such as dimethylformamide with a coupling reagent such as diethyl cyanophosphonate and a base such as diisopropylethylamine at room temperature. A specific example where X is CH₂ is illustrated below.



Alternatively, 2-(1,3-dioxo-1,3-dihydro-isoindol-2-yl)-pentanedioic acid, 35.4, (commercially available) is treated in a solvent such as acetonitrile with triethylamine, 1-hydroxybenzotriazole, 4-methoxybenzylamine, and 1,3-dicyclohexylcarbodiimide. After the reaction is complete, the solvent is removed and the residue is purified by chromatography to generate the desired analog, 35.3, according to a procedure such as that reported in *J. Med. Chem.*, 2003, 46, 3793.

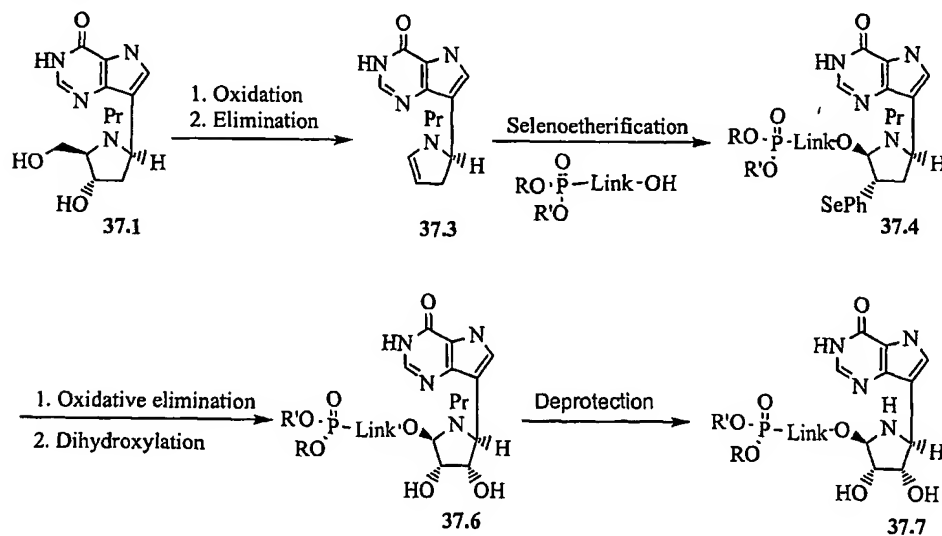
Example 36





The Boc-protected (1*S*)-1-(9-deazaguanin-9-yl)-1,4-dideoxy-1,4-imino-D-ribose, compound, **36.3**, is prepared by stirring the (1*S*)-1-(9-deazaguanin-9-yl)-1,4-dideoxy-1,4-imino-D-ribose (WO 99/19338 and Evans, G. B. et al., *Tetrahedron*, **2000**, *56*, 3053, also reported in Evans, G. B. et al., *J. Med. Chem.* **2003**, *46*, 3412) with BOC anhydride as described in Greene, T., "Protective groups in organic synthesis," Wiley-Interscience, **1999**. Compound, **36.3**, is then treated in a solvent such as tetrahydrofuran or dimethylformamide with a base such as sodium hydride. When bubbling ceases, diethyl phosphonemethyl-triflate (prepared according to *Tetrahedron Lett.*, **1986**, *27*, 1477) is added, yielding the desired phosphonate diester, **36.4**, after deprotection of the BOC group using trifluoroacetic acid (TFA) (which is a compound of Formula 25).

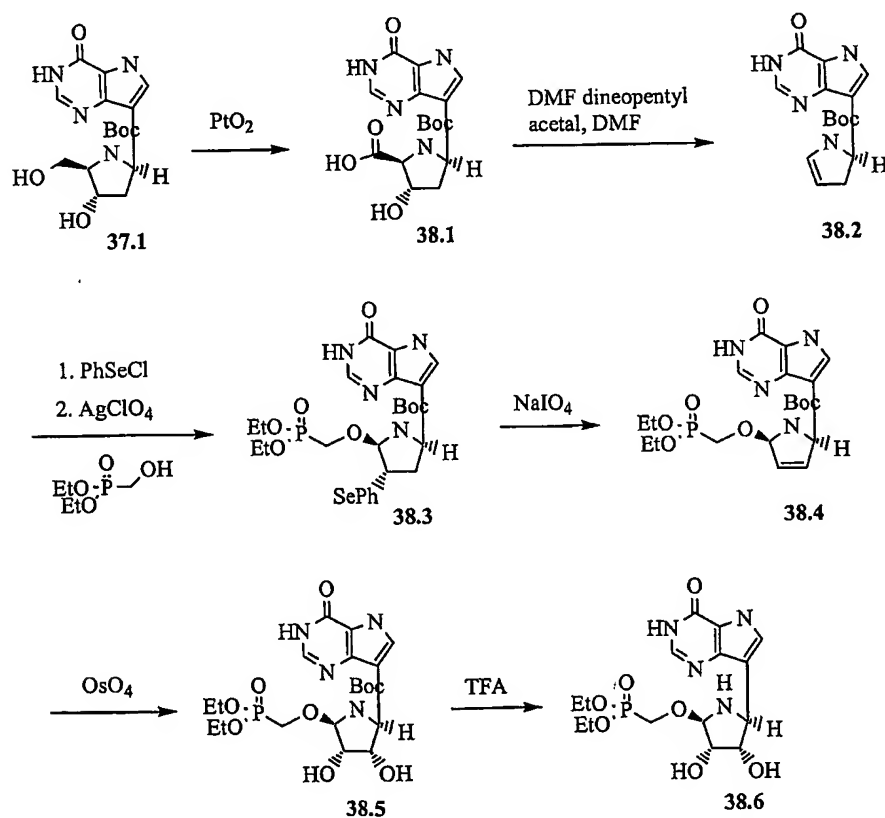
Example 37



Protected compound, **37.1**, ((1*R*)-1-(9-deazahypoxanthin-9-yl)-1,2,4-trideoxy-1,4-imino-D-*erythro*-pentitol, as the hydrochloride salt) is prepared as described in Evans, G. B. et al., *Tetrahedron*, **2000**, *56*, 3053, using di-*t*-butyl

dicarbonate in dichloromethane. Oxidation of the 5'-OH followed by elimination provides glycal, **37.3** (see the procedure of Zemlicka J. et al., *J. Am. Chem. Soc.*, **1972**, *94*, 9, 3213). Selenoetherification provides the protected phosphonate, **37.4** (Kim, C. et al., *J. Org. Chem.*, **1991**, *56*, 2642). Oxidative elimination of the phenylselenide (as described in Kim, C. et al., *J. Org. Chem.*, **1991**, *56*, 2642) followed by stereoselective dihydroxylation provides the desired diol, **37.6**. Finally, the protecting group is removed to provide a compound, **37.7**, compound of Formula 26.

Example 38



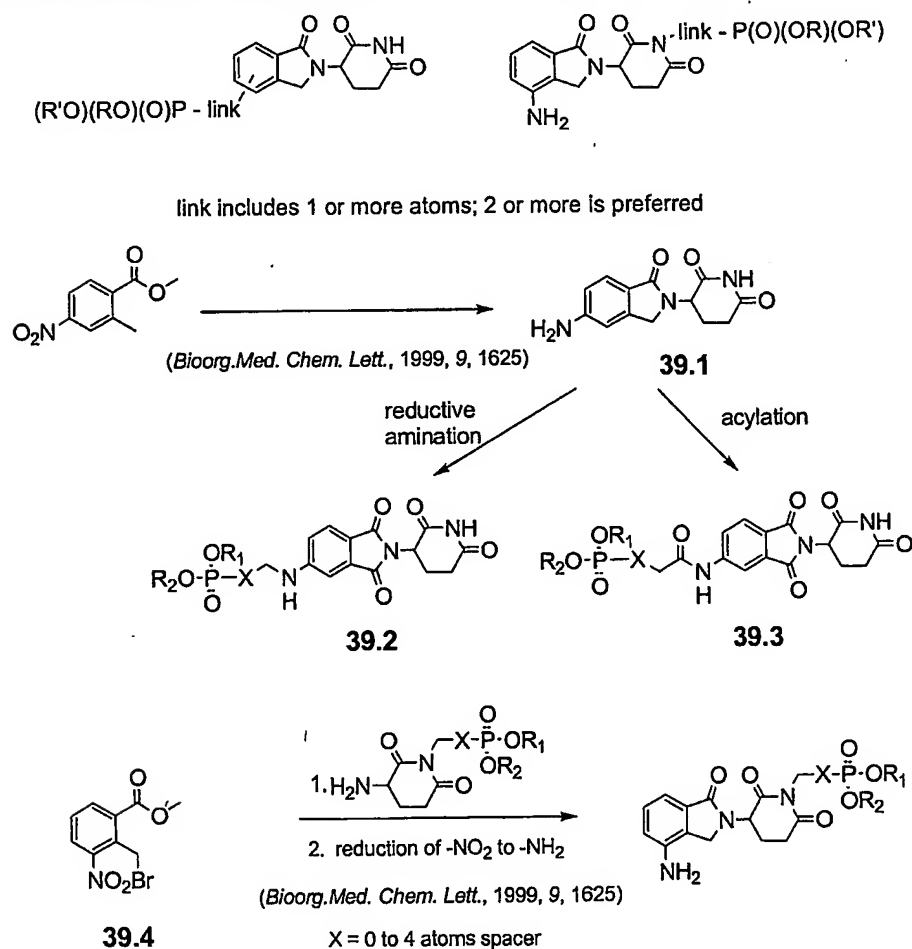
(1R)-1-(9-Deazahypoxanthin-9-yl)-1,2,4-trideoxy-1,4-imino-D-erythro-

pentitol, prepared as the HCl salt as described in Evans, G. B. et al., *Tetrahedron*, **2000**, *56*, 3053, is first protected and then oxidized with PtO_2 to provide carboxylic acid, **38.1**. Decarboxylative elimination is achieved using dimethylformamide dineopentyl acetal in dimethylformamide at high temperature (Zemlicka J. et al., *J. Am. Chem. Soc.*, **1972**, *94*, 9, 3213).

Selenoetherification followed by treatment of the protected glycal with silver perchlorate in the presence of diethyl(hydroxymethyl)phosphonate (Phillion, D. et al., *Tetrahedron Lett.*, 1986, 27, 1477) provides the phosphonate, 38.3 (Kim, C. et al., *J. Org. Chem.*, 1991, 56, 2642). Oxidative elimination of the selenide followed by dihydroxylation using osmium tetroxide provides diol, 38.5. Removal of the amine protecting group, according to the procedure of Greene, T., "Protective groups in organic synthesis," Wiley-Interscience, 1999, provides compound 38.6.

Example 39

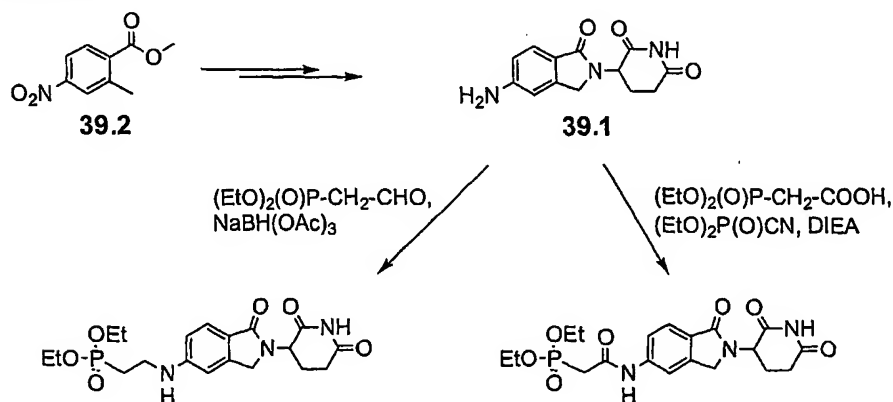
Synthetic methodologies and intermediate compounds that can be used to prepare pro-drugs of analogs of thalidomide are described below.



2-Methyl-4-nitrobenzoic acid methyl ester (commercially available) is converted to 3-(5-amino-1-oxo-1,3-dihydro-isoindol-2-yl)-piperidine-2,6-dione, **39.1**, following the procedures reported in *Bioorg. Med. Chem. Lett.*, 1999, 9, 1625. This amine intermediate is subjected to a reductive amination with diethylphosphonoacetaldehyde (obtained from ozonolysis of diethyl allylphosphonate) in the presence of a reducing agent such as sodium triacetoxyborohydride to generate the desired amine linker analog, **39.2** (*J. Org. Chem.*, 1996, 61, 3849). Alternatively, the amine is acylated with an activated diethylphosphonoacetic acid to provide the desired amide linker compound, **39.3**, according to a procedure such as those reported in *J. Med. Chem.*, 1982, 25, 960 and *J. Med. Chem.*, 1984, 27, 600. The activated diethylphosphonoacetic acid can be obtained by treatment, in a solvent such as dimethylformamide, with a coupling reagent such as diethyl cyanophosphonate and a base such as diisopropylethylamine at room temperature.

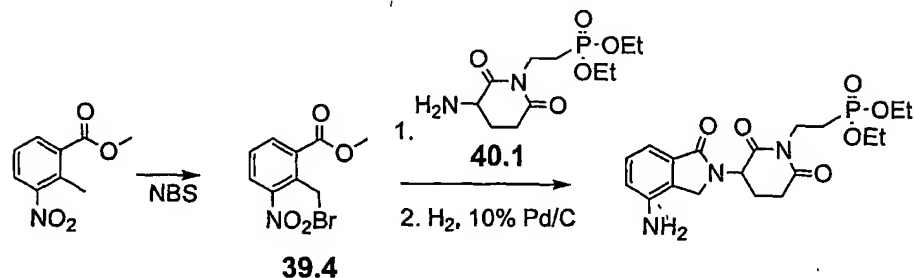
2-Methyl-3-nitrobenzoic acid methyl ester (commercially available) is treated in a solvent such as carbon tetrachloride with N-boromosuccinimide under light to produce 2-bromomethyl-3-nitrobenzoic acid methyl ester, **39.4**. The benzylic bromide is treated in a solvent such as dimethylformamide with [2-(3-amino-2,6-dioxo-piperidin-1-yl)-ethyl]-phosphonic acid diethyl ester (for the preparation of this compound, see Example 40, below) in the presence of a base such as triethylamine. The coupled product is then reduced by hydrogenation (*Bioorg. Med. Chem. Lett.*, 1999, 9, 1625) to afford the desired analog.

Example 40



2-Methyl-4-nitrobenzoic acid methyl ester, **39.2** (commercially available), is converted to 3-(5-amino-1-oxo-1,3-dihydro-isoindol-2-yl)-piperidine-2,6-dione, following the procedures reported in *Bioorg. Med. Chem. Lett.*, 1999, 9, 1625. This amine intermediate is subjected to a reductive amination with diethylphosphonoacetaldehyde (obtained from ozonolysis of diethyl allylphosphonate) in the presence of a reducing agent such as sodium triacetoxyborohydride to generate the desired amine linker analog (*J. Org. Chem.*, 1996, 61, 3849). Alternatively, the amine is acylated with an activated diethylphosphonoacetic acid to provide the desired amide linker compound, according to a procedure such as those reported in *J. Med. Chem.*, 1982, 25, 960 and *J. Med. Chem.*, 1984, 27, 600. The activated diethylphosphonoacetic acid can be obtained by treatment in a solvent such as dimethylformamide with a coupling reagent such as diethyl cyanophosphonate and a base such as diisopropylethylamine at room temperature.

Example 40A



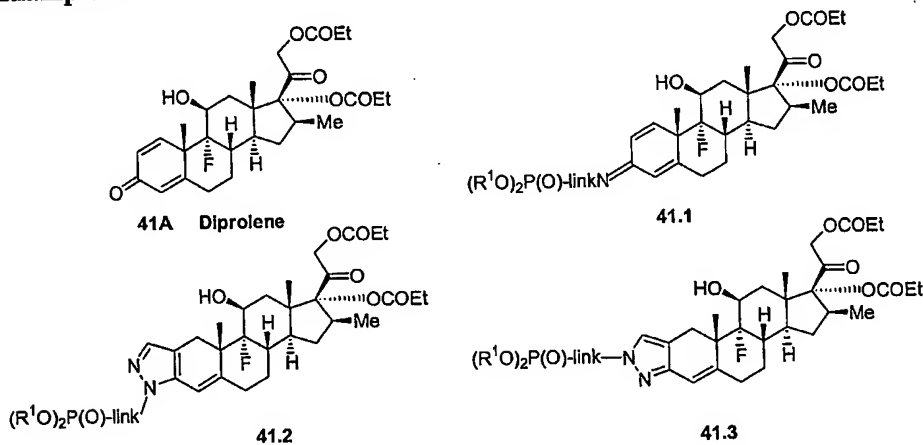
2-Methyl-3-nitrobenzoic acid methyl ester, **39.2** (commercially available) is treated in a solvent such as carbon tetrachloride with N-bromosuccinimide under light to produce 2-bromomethyl-3-nitrobenzoic acid methyl ester, **39.4**. This benzylic bromide is treated in a solvent such as dimethylformamide with [2-(3-amino-2,6-dioxo-piperidin-1-yl)-ethyl]-phosphonic acid diethyl ester (for the preparation of this compound, see below) in the presence of a base such as triethylamine. The coupled product is then reduced by hydrogenation (*Bioorg. Med. Chem. Lett.*, 1999, 9, 1625) to afford the desired analog.

[2-(3-amino-2,6-dioxo-piperidin-1-yl)-ethyl]-phosphonic acid diethyl ester, **40.1**, is obtained according to a procedure such as that reported in *J. Med.*

Chem., 2003, 46, 3793. Accordingly, benzyloxycarbonyl-protected glutaric acid is treated in a solvent such as acetonitrile with triethylamine, 1-hydroxybenzotriazole, diethyl 2-aminoethylphosphonate and 1,3-dicyclohexylcarbodiimide. After the reaction is complete, the solvent is removed and the residue is purified by chromatography to generate the cyclic product, which is subjected to hydrogen in the presence of palladium catalysis to afford the desired intermediate.

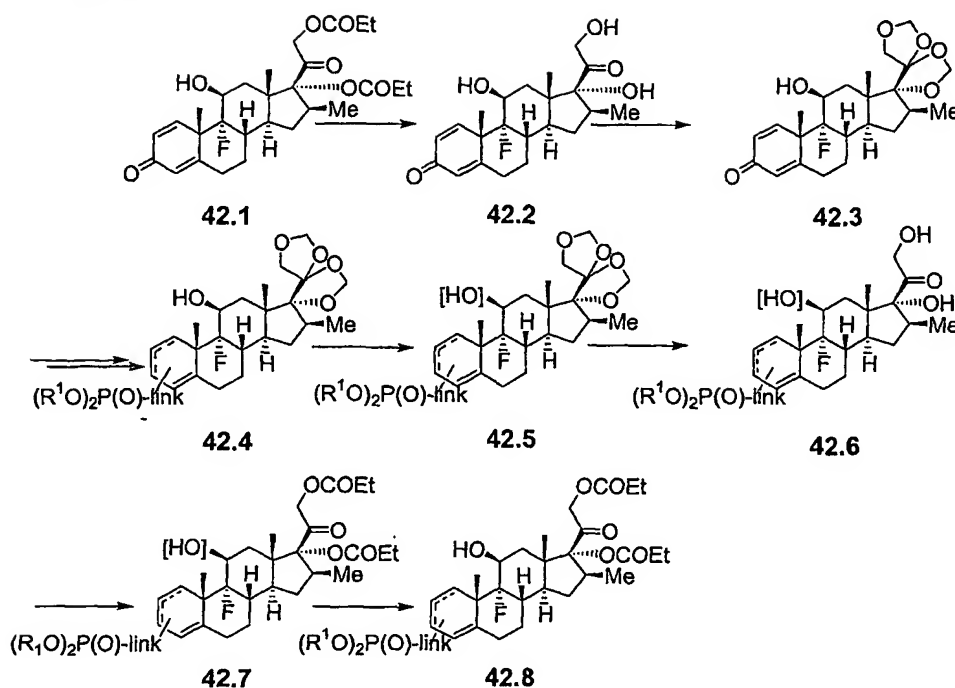
Further manipulations can be performed on the phosphonate moiety prior to the final deprotection. These types of transformations are more extensively described herein.

Example 41



The structures of Diprolene, 41A (German Patent DE 2905674), and the esters, 41.1 – 41.3, are shown below, in which the substituent R^1 is H, alkyl, alkenyl, aryl or aralkyl. The compounds, 41.1 - 41.3, incorporate a phosphonate moiety $(R^1O)_2P(O)$ connected to the nucleus by means of a variable linking group, designated as "link" in the attached structures.

Example 42



A protection-deprotection sequence in which the steroid side-chain is protected as a bis-methylenedioxy (BMD) moiety is shown above. The propionate ester groups of compound, **42.1**, are hydrolyzed, for example by reaction with two molar equivalents of lithium hydroxide in aqueous dimethoxyethane solution at ambient temperature, to give the diol, **42.2**. The product is then reacted with paraformaldehyde and an acid catalyst such as hydrochloric acid, as described in "Protective Groups in Organic Synthesis," by T.W. Greene and P.G.M. Wuts, Wiley, Second Edition 1990, p. 223, to yield the BMD derivative, **42.3**. The phosphonate moiety is then introduced, using the procedures described below, to produce the phosphonate ester, **42.4**. Prior to hydrolysis of the BMD protecting group, the 11-hydroxyl group is protected. The protecting group is selected so that it is stable to the conditions required for removal of the BMD group, and so that it is removable without affecting the subsequently introduced 17, 21-diester moiety.

For example, the 11-hydroxyl group is protected by conversion to the 4-azidobutyrate ester, by reaction with 4-azidobutyryl chloride in pyridine. The 11-azidobutyrate group is then removed from the diester, **42.7**, by reaction with triphenylphosphine, as described in *Bull. Soc. Chem. Jpn.*, 1986, 59, 1296.

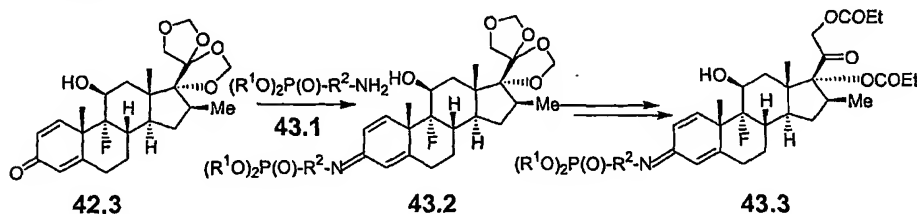
Alternatively, the 11-hydroxyl group is protected by conversion to the 2-(trimethylsilyl)ethyl carbonate, by reaction with 2-(trimethylsilyl)ethyl carbonyl chloride and pyridine. The 2-(trimethylsilyl) carbonate is removed from the diester, **42.7**, by reaction with tetrabutylammonium fluoride in tetrahydrofuran at ambient temperature, as described in *Tet. Lett.*, **1981**, *22*, 969.

Alternatively, the 11-hydroxyl group is protected by conversion to the trichloroacetyl ester, by reaction with trichloroacetyl chloride in dimethylformamide-pyridine. The trichloroacetyl ester is removed by reaction with ethanolic ammonia at ambient temperature, as described in *Coll. Czech. Chem. Commun.*, **1962**, *27*, 2567.

The BMD moiety in the protected product, **42.5**, is then hydrolyzed, for example by treatment with 50% aqueous acetic acid, as described in "Protective Groups in Organic Synthesis," by T.W. Greene and P.G.M. Wuts, Wiley, Second Edition 1990, p. 223, to afford the diol, **42.6**. The diol compound is then acylated, for example by reaction with propionic acid and dicyclohexyl carbodiimide in dimethylformamide at ambient temperature, or by reaction with propionyl chloride and triethylamine in dichloromethane, to produce the dipropionate, **42.7**. Deprotection of the 11-hydroxyl group, as described above, then affords the diester, **42.8**.

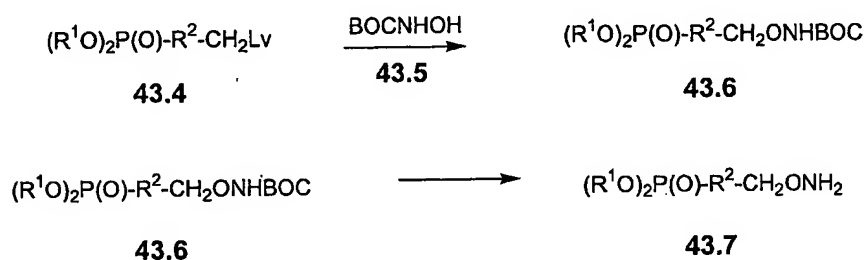
Alternatively, the 20-ketone group is protected as the diethylamine adduct by reaction with titanium tetrakis(diethylamide), as described in Protective Groups in Organic Synthesis, by T.W. Greene and P.G.M Wuts, Wiley, Second Edition 1990, p. 219.

Example 43



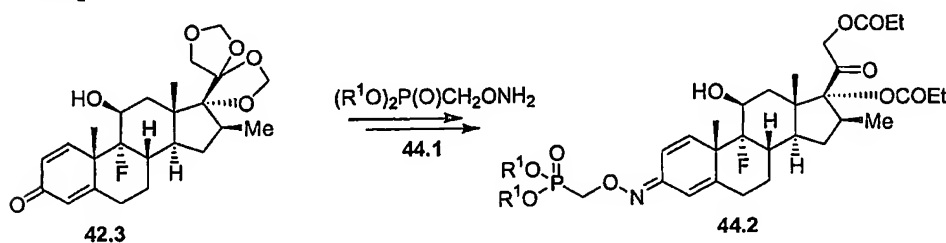
The preparation of phosphonates in which the phosphonate is attached by means of an imino or iminoxy group and a variable carbon chain is illustrated above. In this procedure, the BMD-protected derivative, **42.3**, is reacted with an amine or hydroxylamine, **43.1**, in which R^2 is an alkyl, alkenyl, cycloalkyl or

cycloalkenyl group, optionally incorporating a heteroatom O, S or N, or a functional group such as an amide, ester, oxime, sulfoxide or sulfone etc, or an optionally substituted aryl, heteroaryl or aralkyl group, optionally incorporating a heteroatom O, S or N, to afford the imine or iminoxy product, 43.2. The reaction is conducted between equimolar amounts of the reactants in an aprotic solvent such as pyridine or xylene, or in an alcoholic solvent such as ethanol, optionally in the presence of an acid catalyst, to give the imine or oxime. The preparation of oximes of steroidal 3-ketones is described in *Anal. Bioch.*, 1978, 86, 133 and in *J. Mass. Spectrom.*, 1995, 30, 497. The BMD-protected compound, 43.2, is then converted, as described above, into the diester, 43.3.



The preparation of hydroxylamine ethers incorporating a phosphonate group is shown above. In this procedure, a phosphonate, 43.4, in which Lv is a leaving group such as bromo or trifluoromethylsulfonyloxy, is reacted with BOC-hydroxylamine, 43.5 (Aldrich), to produce the ether, 43.6. The reaction is conducted between equimolar amounts of the reactants in a polar solvent such as dimethylformamide or tetrahydrofuran, in the presence of a base such as potassium hydroxide or dimethylaminopyridine, to give the product, 43.6. Deprotection, for example by treatment with trifluoroacetic acid, then gives the hydroxylamine ether, 43.7.

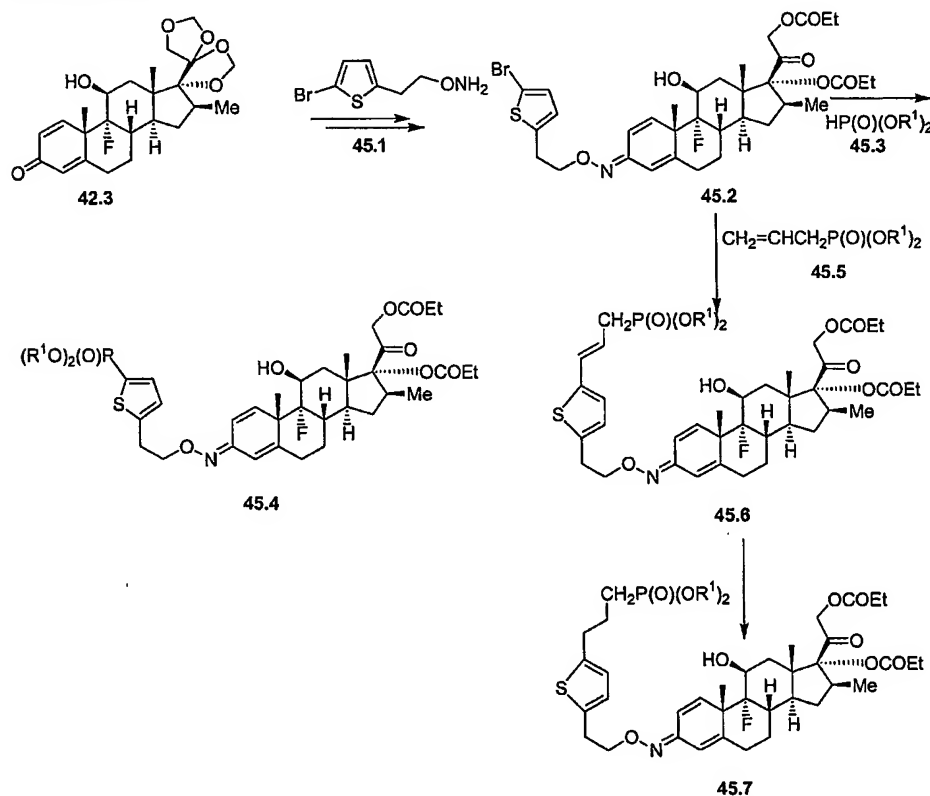
Example 44



The preparation of phosphonates in which the phosphonate is attached by means of an iminoxy group is illustrated above. In this procedure, the substrate, **42.3**, is reacted with a dialkyl phosphonomethyl hydroxylamine, **44.1**, prepared as described above from a dialkyl trifluoromethylsulfonyloxymethyl phosphonate (*Tetrahedron Lett.*, 1986, 27, 1477) and BOC-hydroxylamine, to afford, after deprotection and side chain acylation, the oxime ether, **44.2**. The oxime forming reaction is performed at ambient temperature in pyridine solution between equimolar amounts of the reactants.

Using the above procedures, but employing, in place of the oxime ether, **44.1**, different oxime ethers, **43.7**, the corresponding products, **43.3** are obtained.

Example 45



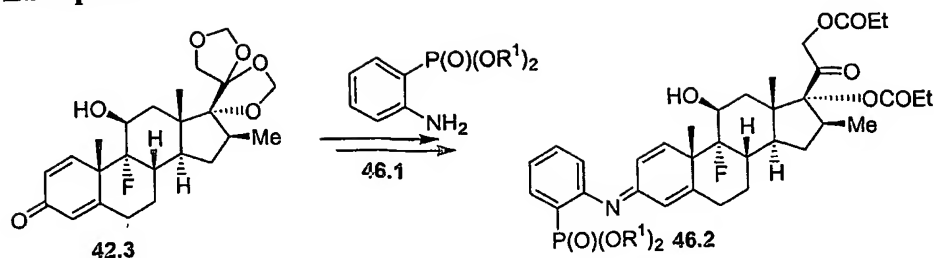
The preparation of phosphonates incorporating an iminoxy group, by means of the reaction between the substrate the substrate, **42.3**, and O-2-(5-bromo-2-thienyl)ethoxyhydroxylamine, **45.1**, prepared as described above from 2-(5-bromo-2-thienyl)ethyl bromide (*J. Chem. Soc., Perkin Trans. Phys. Org. Chem.*, 1975, 821), is illustrated above. The resultant oxime ether is converted,

by deprotection and side chain acylation, into the compound, 45.2, which is then reacted, in the presence of a palladium catalyst, with a dialkyl phosphite, 45.3, to afford the phosphonate, 45.4. The preparation of arylphosphonates by means of a coupling reaction between aryl bromides and dialkyl phosphites is described in *J. Med. Chem.*, 1992, 35, 1371. The reaction is performed in an inert solvent such as toluene, in the presence of a base such as triethylamine and a catalytic amount of tetrakis(triphenylphosphine)palladium(0).

Alternatively, the bromo-substituted product, 45.2, is coupled, in a palladium-catalyzed Heck reaction, with a dialkyl propenyl phosphonate, 45.5, (Acros) to give the unsaturated phosphonate, 45.6. The coupling of aryl halides with olefins by means of the Heck reaction is described, for example, in "Advanced Organic Chemistry," by F. A. Carey and R. J. Sundberg, Plenum, 2001, p. 503ff and in *Acc. Chem. Res.*, 1979, 12, 146. The aryl bromide and the olefin are coupled in a polar solvent such as dimethylformamide or dioxan, in the presence of a palladium(0) catalyst such as tetrakis(triphenylphosphine)-palladium(0) or palladium(II) catalyst such as palladium(II) acetate, and optionally in the presence of a base such as triethylamine or potassium carbonate.

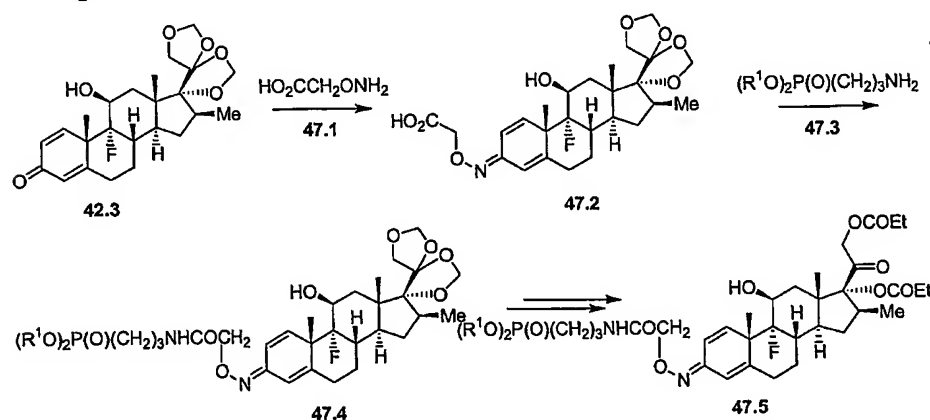
Optionally, the styrenoid double bond present in the product, 45.6, is reduced, for example by reaction with diimide, to produce the saturated analog, 45.7. The reduction of olefinic bonds is described in "Comprehensive Organic Transformations," by R. C. Larock, VCH, 1989, p. 6ff. The transformation is effected by means of catalytic hydrogenation, for example using a palladium on carbon catalyst and hydrogen, or a hydrogen donor, or by the use of diimide or diborane.

Using the above procedures, but employing, in place of the bromothieryl reagent, 45.1, different bromo-substituted aryl or heteroaryl alkoxy hydroxylamines, and/or different dialkyl alkenyl phosphonates, the products analogous to the compounds 45.4, 45.6 and 45.7 are obtained.

Example 46

The preparation of phosphonates in which the phosphonate is attached by means of an imino group is illustrated above. In this procedure, the substrate, 42.3, which is reacted with a dialkyl 2-aminophenyl phosphonate, 46.1 (Aurora), to give, after deprotection and side chain acylation, the imine product, 46.2. The reaction is conducted in a hydrocarbon solvent such as toluene or xylene, at reflux temperature, in the presence of a basic catalyst such as sodium methoxide, or an acid catalyst such as p-toluenesulfonic acid, under azeotropic conditions, to give the product, 46.2.

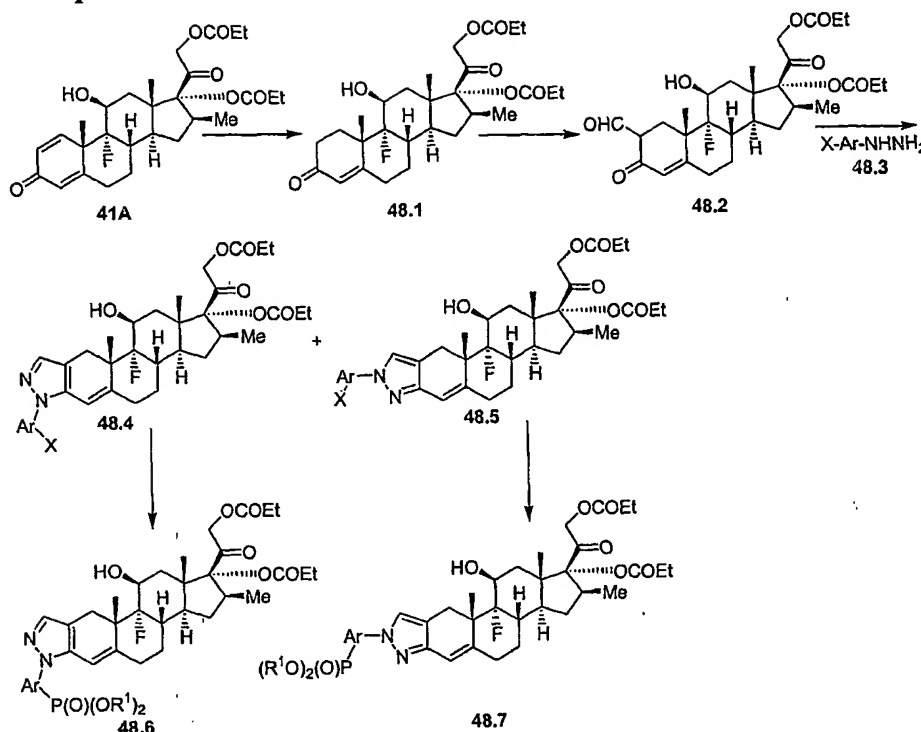
Using the above procedures, but employing, in place of the 2-amino-phenyl phosphonate, 46.1, different amino-substituted aryl or heteroaryl phosphonates, products analogous to 46.2 are obtained.

Example 47

An alternative method for the preparation of phosphonates in which the phosphonate is attached by means of an oximino group is illustrated above. In this procedure, the dienone, 42.3, is reacted with O-(carboxymethyl)hydroxylamine, 47.1 (Interchim), to yield, after deprotection and side chain acylation, the

oxime, 47.2. The reaction of steroidal 1,4-dien-3-ones with hydroxylamine is described in *J. Steroid Bioch.*, 1976, 7, 795. The reaction is performed between equimolar amounts of the reactants in a polar organic solvent such as pyridine or methanol, optionally in the presence of acetic acid or sodium acetate. The oxime, 47.2, is then reacted with a dialkyl 3-hydroxyphenyl phosphonate, 47.3 (Epsilon), in a Mitsunobu reaction, to yield the substituted oxime, 47.4. The preparation of aromatic ethers and thioethers by means of the Mitsunobu reaction is described, for example, in "Comprehensive Organic Transformations," by R. C. Larock, VCH, 1989, p. 448, and in "Advanced Organic Chemistry," Part B, by F.A. Carey and R. J. Sundberg, Plenum, 2001, p. 153-4 and in *Org. React.*, 1992, 42, 335. The phenol and the hydroxy or mercapto component are reacted together in an aprotic solvent such as, for example, tetrahydrofuran, in the presence of a dialkyl azodicarboxylate and a triarylphosphine, to afford the ether or thioether products. The procedure is also described in *Org. React.*, 1992, 42, 335-656. The product, 47.4, is then transformed, by deprotection and acylation, into the diester, 47.5.

Using the above procedures, but employing, in place of the phosphonate, 47.3, different dialkyl hydroxy-substituted aryl or heteroaryl phosphonates, the products analogous to 47.5 are obtained.

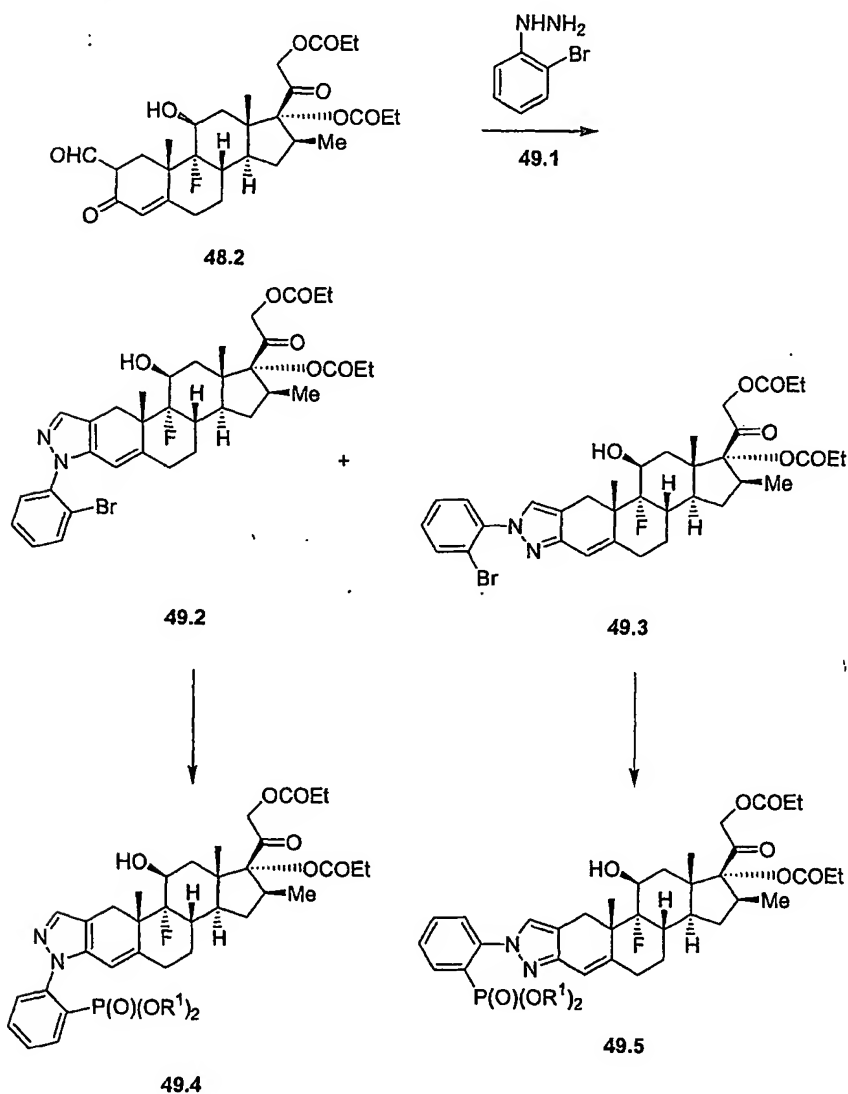
Example 48

The preparation of the phosphonate esters in which the phosphonate group is attached to the 1' or 2' position of the pyrazole ring, by means of an aromatic or heteroaromatic group, a heteroatom and a variable carbon chain is illustrated above. In this procedure, Dipropylene, 41A, is reduced to afford the 1,2-dihydro product, 48.1. The catalytic hydrogenation reaction is effected by the use of tris(triphenylphosphine)rhodium (I) chloride, for example as described in *J. Med. Chem.*, 2001, 44, 602. The product is then reacted with ethyl formate and a base such as sodium hydride, in an inert solvent such as toluene or dimethylformamide, as described in Australian Patent Application 275950409, to afford the 2-formyl product, 48.2.

Optionally, the substrate, 41A, is protected, for example as described in Example 42, above, prior to the formylation reaction, as described in *J. Am. Chem. Soc.*, 1964, 86, 1520. The 2-formyl product is then reacted with an aryl or heteroaryl hydrazine, 48.3, in which the substituent X is either a phosphonate group or a group which is subsequently transformed into a phosphonate-containing substituent. For example, X is dialkylphosphono, bromo, hydroxy, amino, carboxyl and the like. The reaction yields the isomeric 2'- and 1'-aryl pyrazoles, 48.4 and 48.5. The ring-forming reaction is performed between

equimolar amounts of the reactants in an acidic solvent such as acetic acid, as described in *J. Am. Chem. Soc.*, 1964, 86, 1520. The pyrazoles, 48.4 and 48.5 are then transformed, for example by the procedures described in Examples 49-50, respectively into the phosphonates, 48.6 and 48.7.

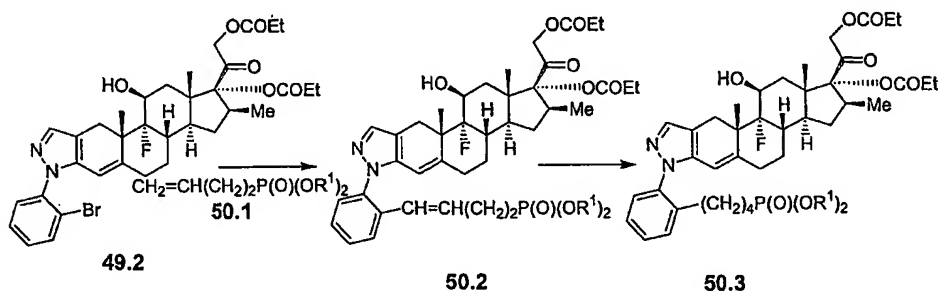
Example 49



The preparation of phosphonates in which the phosphonate is attached by means of a phenyl group is illustrated above. In this sequence, the ketoaldehyde, 48.2, is reacted, as described above, with 2-bromophenylhydrazine, 49.1 (Fluka), to give the isomeric pyrazole products, 49.2 and 49.3. The products are then

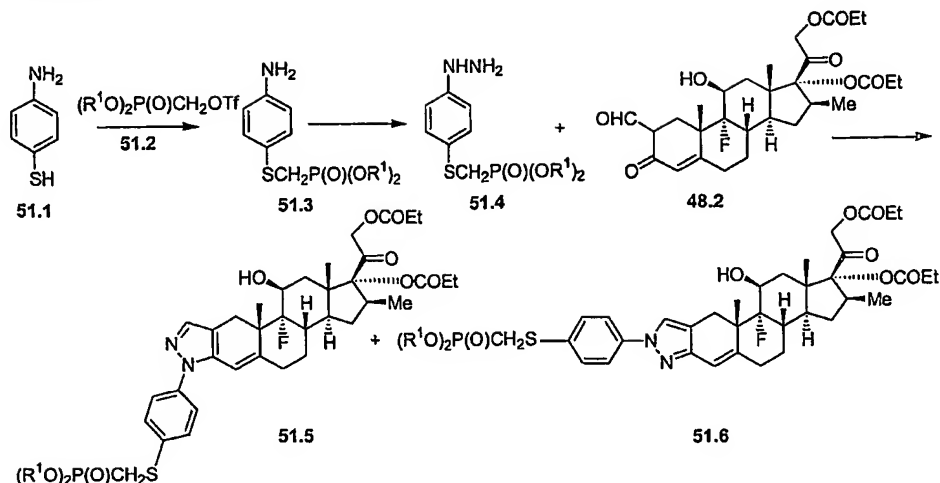
reacted, as described herein, with a dialkyl phosphite $\text{HP(O)(OR}^1)_2$ and a palladium catalyst, to afford respectively the phosphonates, 49.4 and 49.5. Using the above procedures, but employing, in place of 2-bromophenyl hydrazine, different bromoaryl or bromoheteroaryl hydrazines, 48.3, the products, 48.6 and 48.7 are obtained.

Example 50



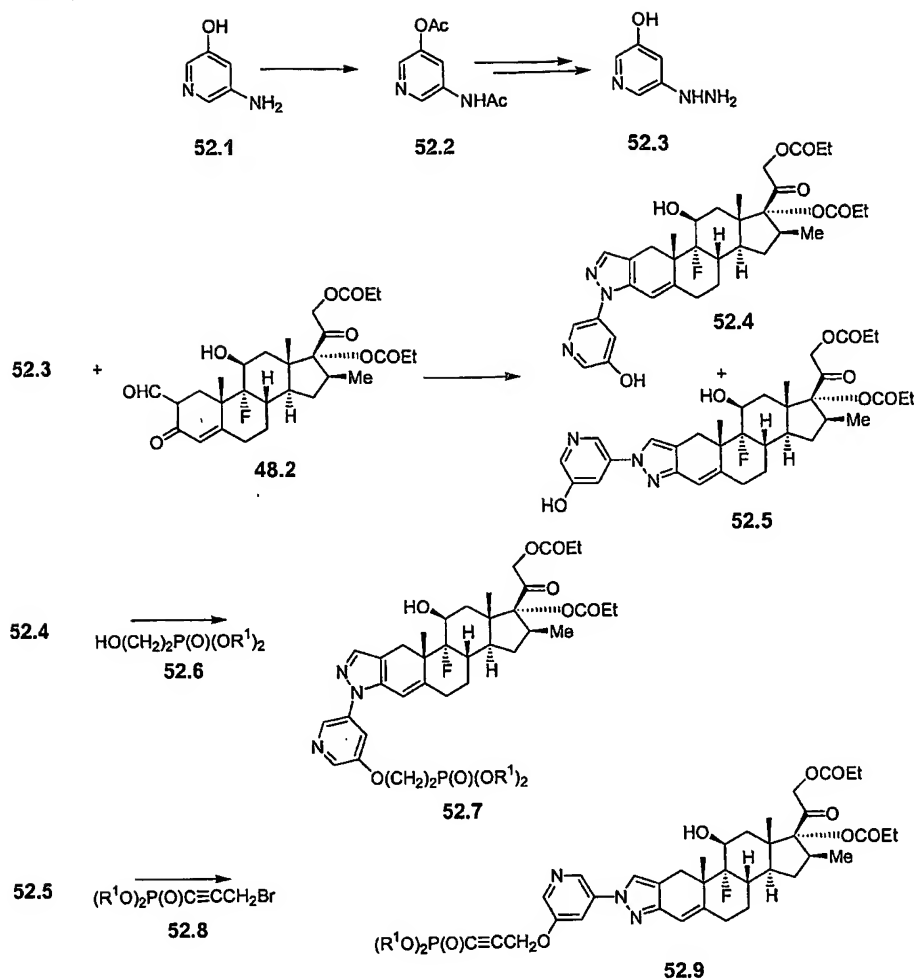
The preparation of phosphonates in which the phosphonate is attached by means of an aromatic or heteroaromatic group and a saturated or unsaturated alkyl chain is illustrated above. In this procedure, the bromophenyl-substituted pyrazole, 49.2, is coupled in a Heck reaction, as described above, with, for example a dialkyl butenyl phosphonate, 50.1, (*Org. Lett.*, 2001, 3, 217) to give the unsaturated phosphonate product, 50.2. Optionally, the product is reduced, as described above, to give the saturated analog, 50.3. Application of the above procedures to the isomeric bromophenyl pyrazole, 49.3, affords the products isomeric with, 50.2 and 50.3. Using the above procedures, but employing, in place of the phosphonate, 50.1, different dialkyl alkenyl phosphonates, and/or different bromoaryl or heteroaryl pyrazoles, 48.4 or 48.5, the products analogous to 50.2 and 50.3 are obtained.

Example 51



The preparation of phosphonates, **51.5** and **51.6**, in which the phosphonate is attached by means of an aryl or heteroaryl group and an alkoxy chain is illustrated above. In this procedure, 4-Aminothiophenol, **51.1**, is reacted in dimethylformamide solution at ambient temperature with a dialkyl trifluoromethanesulfonyloxymethyl phosphonate, **51.2** (*Tetrahedron Lett.*, 1986, 27, 1477), and potassium carbonate to give the thioether, **51.3**. The product is then converted into the corresponding hydrazine, **51.4**, by means of a diazotization reaction in aqueous ethanolic hydrochloric acid, followed by reduction of the diazonium chloride with tin(II) chloride, as described in *J. Med. Chem.*, 2001, 44, 4031. The hydrazine is then reacted, as described above, with the ketoaldehyde, **48.2**, to form the isomeric pyrazoles, **51.5** and **51.6**.

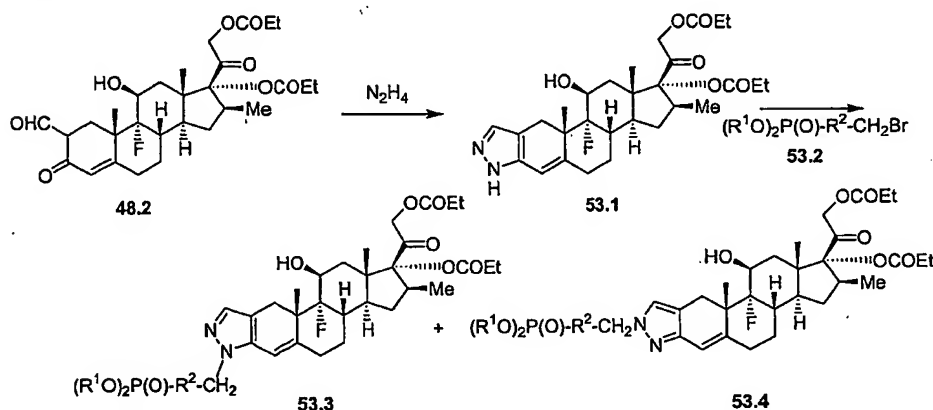
Using the above procedures, but employing, in place of the triflate, **51.2**, different dialkylphosphono alkyl bromides or triflates, and/or different aromatic or heteroaromatic mercapto or hydroxyamines, the products analogous to **51.5** and **51.6** are obtained.

Example 52

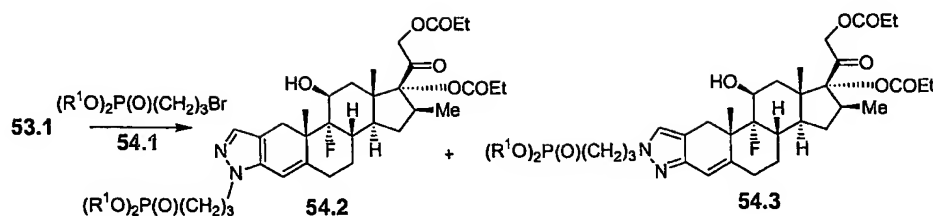
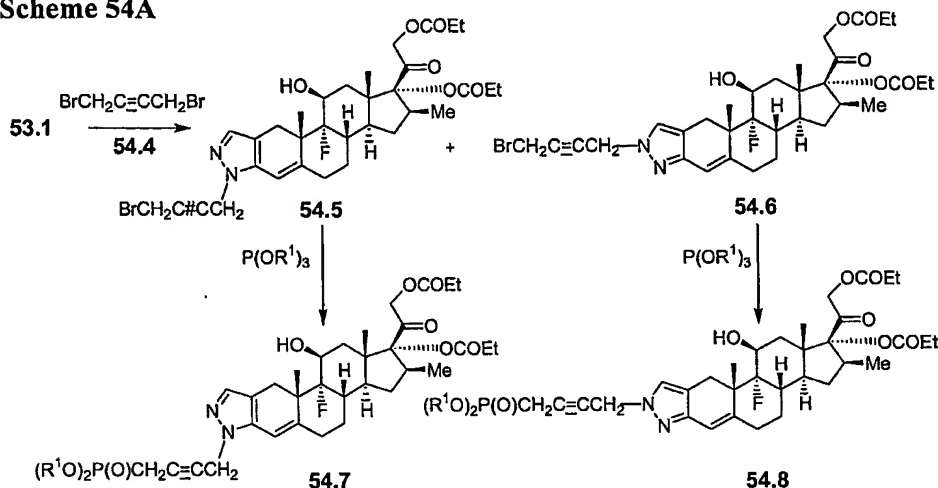
The preparation of phosphonates in which the phosphonate is attached by means of a pyridyl group a heteroatom and a variable carbon chain is illustrated above. In this procedure, 3-amino-5-hydroxypyridine, 52.1, is converted, by reaction with acetic anhydride, into the diacetyl analog, 52.2. The product is then transformed by diazotization and reduction, as described above, into the hydrazine, 52.3. The hydrazine, 52.3, is then reacted with the ketoaldehyde, 48.2, to give the isomeric pyrazoles, 52.4 and 52.5. The 2'-pyridyl product, 52.4, is reacted in a Mitsunobu reaction, as described above, with a dialkyl hydroxyethyl phosphonate, 52.6 (*Zh. Obshchei. Khim.*, 1973, 43, 2364), to afford the ether, 52.7. Application of this procedure to the isomeric phenol, 52.5, affords the product isomeric to compound, 52.7.

Alternatively, the isomeric phenol, **52.5**, is reacted, in dimethylformamide solution at about 80°, with one molar equivalent of a dialkyl bromopropynyl phosphonate, **52.8** (*Bioorg. Med. Chem. Lett.*, 1994, 4, 273), and cesium carbonate, to prepare the phosphonate, **52.9**. Application of this procedure to the isomeric phenol, **52.4**, affords the product isomeric with compound, **52.4**. Using the above procedures, but employing, in place of the carbinol, **52.6**, or the bromide, **52.8**, different thiols, alcohols or bromides, and/or different phenols, **48.4** or **48.5**, in which X is OH, the corresponding products analogous to **52.7** and **52.9** are obtained.

Example 53



The preparation of the phosphonate esters in which the phosphonate group is attached by means of a variable carbon linkage is illustrated above. In this procedure, the ketoaldehyde, **48.2**, is reacted with hydrazine, to afford the pyrazole derivative, **53.1**. The reaction of steroidal 2-formyl-3-ketones with hydrazine is described in *J. Am. Chem. Soc.*, 1964, 86, 1520. The reaction is performed in acetic acid at ambient temperature. The resulting pyrazole is then reacted with a dialkyl bromomethyl phosphonate, **53.2**, in which R^2 is as defined above, to produce the isomeric 2' and 1' alkylation products, **53.3** and **53.4**, respectively. The alkylation of substituted pyrazoles is described, for example, in "Heterocyclic Chemistry," by T. L. Gilchrist, Longman, 1992, p. 309.

Example 54**Scheme 54A**

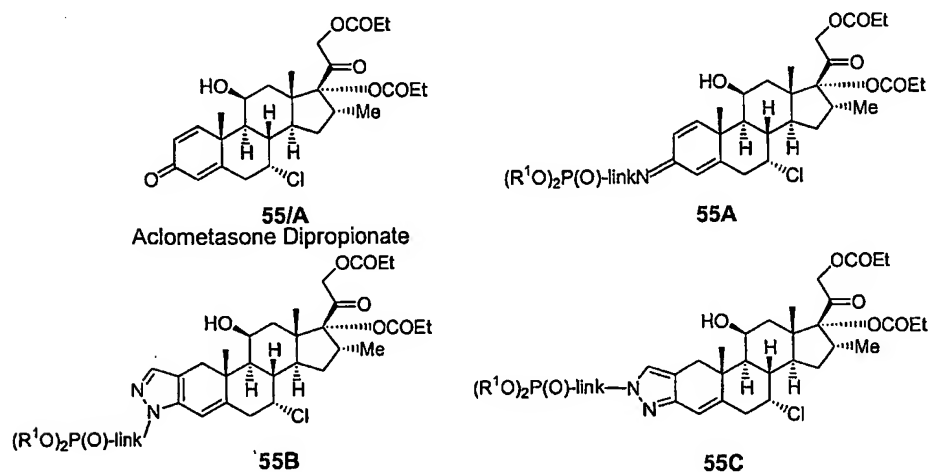
Pyrazole, **53.1**, is reacted, in dimethylformamide solution at ca. 90°, with a dialkyl bromopropyl phosphonate, **54.1** (Aldrich), and a base such as dimethylaminopyridine or lithium hexamethyldisilazide, to yield the isomeric alkylation products, **54.2** and **54.3**.

As shown in Scheme 54A, the pyrazole, **53.1**, is reacted in dimethylformamide solution at ambient temperature with one molar equivalent of 1,4-dibromobut-2-yne, **54.4** (Narchem), and potassium carbonate, to afford the alkylation products, **54.5** and **54.6**. The products are then heated at 120° with a trialkyl phosphite in an Arbuzov reaction, to yield the phosphonates, **54.7** and **54.8**. The Arbuzov reaction is described in *Handb. Organophosphorus Chem.*, 1992, 115-72. Using the above procedures, but employing, in place of the dibromide, **54.4**, different alkyl, alkenyl or alkynyl dibromides, the products analogous to **54.7** and **54.8** are obtained.

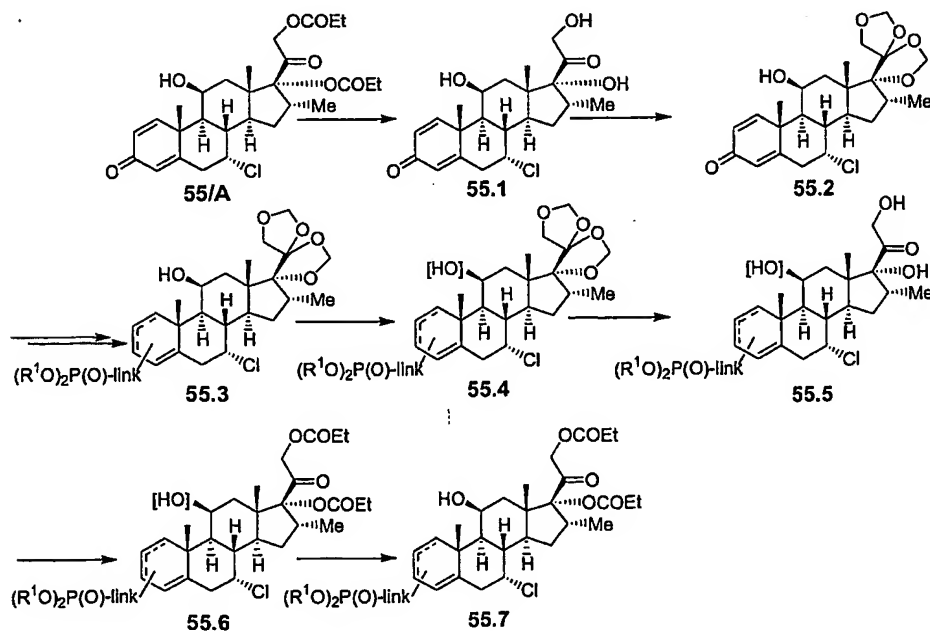
Example 55

The structures of Aclometasone dipropionate, **55/A** (*J. Med. Chem.*, 1980, 23, 430; U.S. Patent No. 4,124,707), and the esters, **55A** – **55C**, are shown

below, in which the substituent R^1 is H, alkyl, alkenyl, aryl or aralkyl. The compounds, **55A** – **55C**, incorporate a phosphonate moiety $(R^1O)_2P(O)$ connected to the nucleus by means of a variable linking group, designated as "link" in the attached structures.



Scheme 55A



The steroid side-chain is protected as a bis-methylenedioxy (BMD) moiety, as illustrated in Scheme 55A. In this sequence, the propionate esters are hydrolyzed, for example by reaction with two molar equivalents of lithium hydroxide in aqueous dimethoxyethane solution at ambient temperature, to give

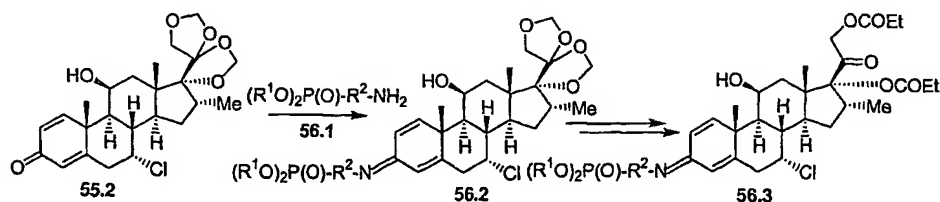
the diol, **55.1**. The product is then reacted with paraformaldehyde and an acid catalyst such as hydrochloric acid, as described in "Protective Groups in Organic Synthesis," by T.W. Greene and P.G.M. Wuts, Wiley, Second Edition 1990, p. 223, to yield the BMD derivative, **55.2**. The phosphonate moiety is then introduced, using the procedures described below, to produce the phosphonate ester, **55.3**. Prior to hydrolysis of the BMD protecting group, the 11-hydroxyl group is protected. The protecting group is selected so that it is stable to the conditions required for removal of the BMD group, and so that it is removable without affecting the subsequently introduced 17, 21-diester moiety. For example, the 11-hydroxyl group is protected by conversion to the 4-azido-butyrate ester, by reaction with 4-azidobutyryl chloride in pyridine. The 11-azidobutyrate group is then removed from the diester, **55.6**, by reaction with triphenylphosphine, as described in *Bull. Soc. Chem. Jpn.*, **1986**, 59, 1296. Alternatively, the 11-hydroxyl group is protected by conversion to the 2-(trimethylsilyl)ethyl carbonate, by reaction with 2-(trimethylsilyl)ethyl carbonyl chloride and pyridine. The 2-(trimethylsilyl) carbonate is removed from the diester, **55.6**, by reaction with tetrabutylammonium fluoride in tetrahydrofuran at ambient temperature, as described in *Tet. Lett.*, **1981**, 22, 969.

Alternatively, the 11-hydroxyl group is protected by conversion to the trichloroacetyl ester, by reaction with trichloroacetyl chloride in dimethylformamide/pyridine. The trichloroacetyl ester is removed by reaction with ethanolic ammonia at ambient temperature, as described in *Coll. Czech. Chem. Commun.*, **1962**, 27, 2567. The BMD moiety in the protected product, **55.4**, is then hydrolyzed, for example by treatment with 50% aqueous acetic acid, as described in "Protective Groups in Organic Synthesis," by T.W. Greene and P.G.M. Wuts, Wiley, Second Edition 1990, p. 223, to afford the diol, **55.5**. The diol, **55.5**, is acylated, for example by reaction with propionic acid and dicyclohexyl carbodiimide in dimethylformamide at ambient temperature, or by reaction with propionyl chloride and triethylamine in dichloromethane, to produce the dipropionate, **55.6**. Deprotection of the 11-hydroxyl group, as described above, then affords the diester, **55.7**.

Alternatively, the 20-ketone group is protected as the diethylamine adduct by reaction with titanium tetrakis(diethylamide), as described in

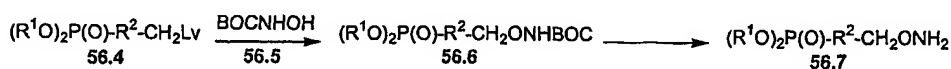
"Protective Groups in Organic Synthesis," by T.W. Greene and P.G.M Wuts, Wiley, Second Edition 1990, p. 219.

Example 56



The preparation of phosphonates in which the phosphonate is attached by means of an imino or iminoxy group and a variable carbon chain is illustrated above. In this procedure, the BMD-protected derivative, 55.2, is reacted with an amine or hydroxylamine, 56.1, in which R^2 is an alkyl, alkenyl, cycloalkyl or cycloalkenyl group, optionally incorporating a heteroatom O, S or N, or a functional group such as an amide, ester, oxime, sulfoxide or sulfone etc, or an optionally substituted aryl, heteroaryl or aralkyl group, optionally incorporating a heteroatom O, S or N, to afford the imine or iminoxy product, 56.2. The reaction is conducted between equimolar amounts of the reactants in an aprotic solvent such as pyridine or xylene, or in an alcoholic solvent such as ethanol, optionally in the presence of an acid catalyst to give the imine or oxime. The preparation of oximes of steroidal 3-ketones is described in *Anal. Bioch.*, 1978, 86, 133 and in *J. Mass. Spectrom.*, 1995, 30, 497. The BMD-protected side-chain compound, 56.2, is then converted into the diester, 56.3.

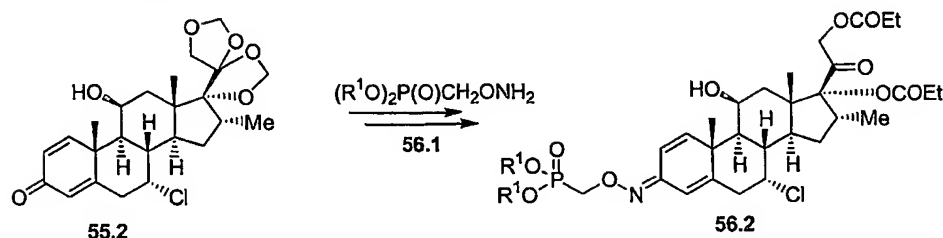
Example 56A



Also illustrated is the preparation of hydroxylamine ethers incorporating a phosphonate group. A phosphonate, 56.4, in which Lv is a leaving group such as bromo or trifluoromethylsulfonyloxy, is reacted with BOC-hydroxylamine, 56.5 (Aldrich), to produce the ether, 56.6. The reaction is conducted between equimolar amounts of the reactants in a polar solvent such as dimethylformamide or tetrahydrofuran, in the presence of a base such as potassium hydroxide or dimethylaminopyridine, to give the product, 56.6. Deprotection,

for example by treatment with trifluoroacetic acid, then gives the hydroxylamine ether, **56.7**.

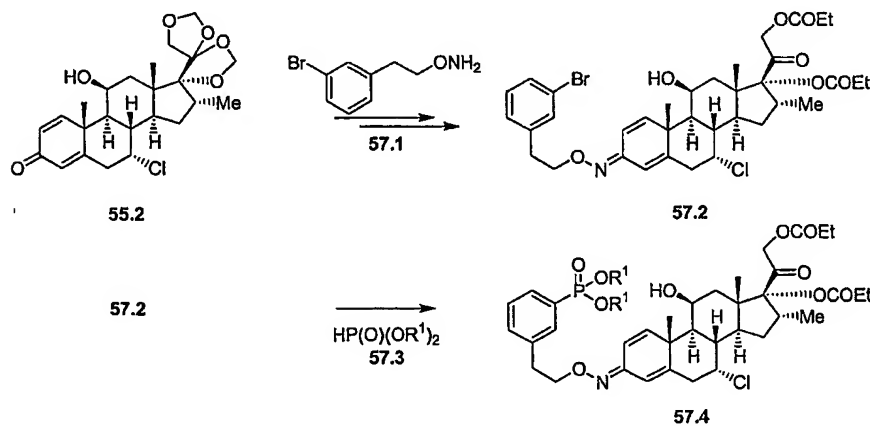
Example 56B



The preparation of phosphonates in which the phosphonate is attached by means of an iminoxy group is illustrated above. In this procedure, the substrate, **55.2**, is reacted with a dialkyl phosphonomethyl hydroxylamine, **56.1**, prepared as described above from a dialkyl trifluoromethylsulfonyloxymethyl phosphonate (*Tetrahedron Lett.*, 1986, 27, 1477) and BOC-hydroxylamine, to afford, after deprotection and side chain acylation, the oxime ether, **56.2**. The oxime forming reaction is performed at ambient temperature in pyridine solution between equimolar amounts of the reactants.

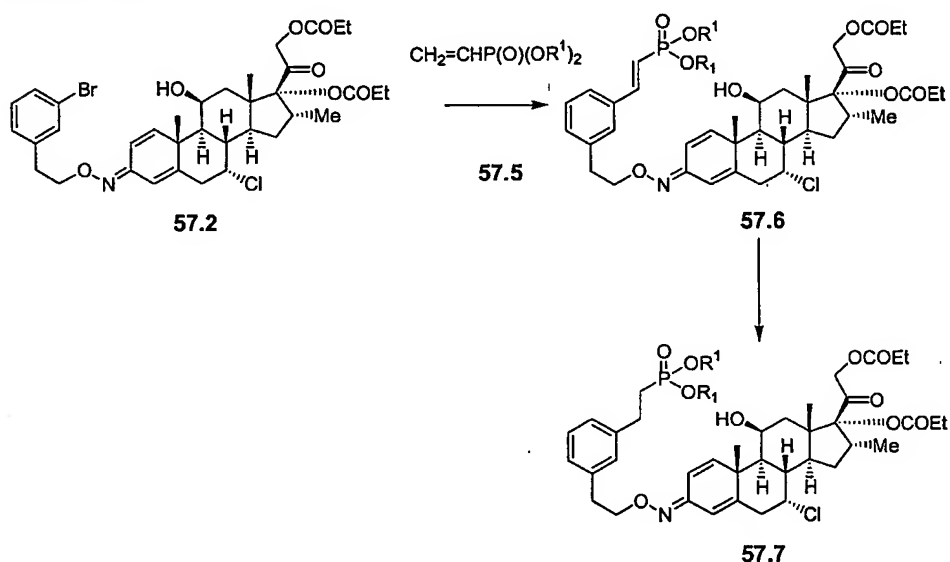
Using the above procedures, but employing, in place of the oxime ether, **56.1**, different oxime ethers, **56.7**, the corresponding products, **56.2** are obtained.

Example 57



The preparation of phosphonates incorporating an iminoxy group, by means of the reaction between the substrate, **55.2**, and O-2-(3-bromophenyl)-ethylhydroxylamine, **57.1**, prepared as described above from compound, **55.1**,

and 2-(3-bromophenyl)ethyl bromide, respectively. The resultant oxime ether is converted, by deprotection and side chain acylation, into the compound, **57.2**, which is then reacted, in the presence of a palladium catalyst, with a dialkyl phosphite, **57.3**, to afford the phosphonate, **57.4**. The preparation of arylphosphonates by means of a coupling reaction between aryl bromides and dialkyl phosphites is described in *J. Med. Chem.*, 1992, 35, 1371. The reaction is performed in an inert solvent such as toluene, in the presence of a base such as triethylamine and a catalytic amount of tetrakis(triphenylphosphine)-palladium(0).

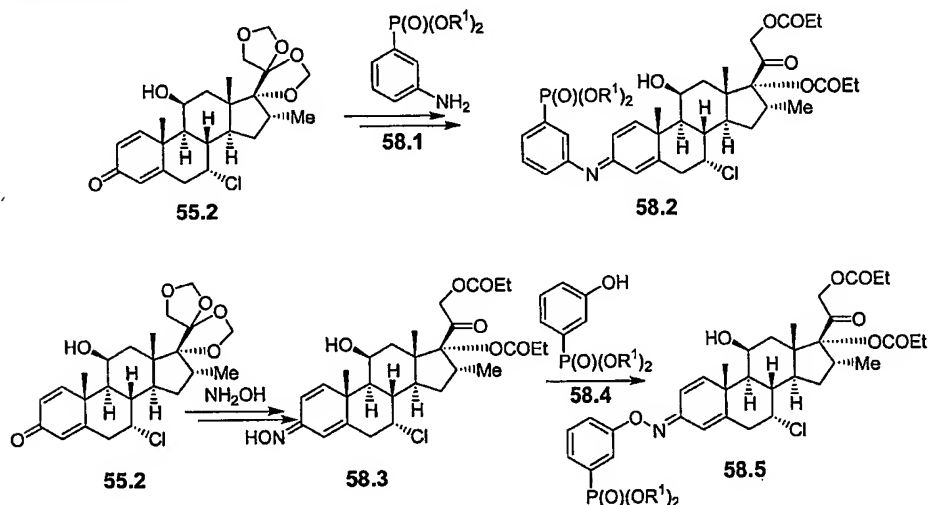


Alternatively, the bromo-substituted product, **57.2**, is coupled, in a palladium-catalyzed Heck reaction, with a dialkyl vinyl phosphonate, **57.5** (Aldrich), to give the unsaturated phosphonate, **57.6**. The coupling of aryl halides with olefins by means of the Heck reaction is described, for example, in "Advanced Organic Chemistry," by F. A. Carey and R. J. Sundberg, Plenum, 2001, p. 503ff and in *Acc. Chem. Res.*, 1979, 12, 146. The aryl bromide and the olefin are coupled in a polar solvent such as dimethylformamide or dioxan, in the presence of a palladium(0) catalyst such as tetrakis(triphenylphosphine)-palladium(0) or a palladium(II) catalyst such as palladium(II) acetate, and optionally in the presence of a base such as triethylamine or potassium carbonate. Optionally, the styrenoid double bond present in the product, **57.6**, is reduced, for example by reaction with diimide, to produce the saturated analog, **57.7**. The reduction of olefinic bonds is described in "Comprehensive Organic

Transformations," by R. C. Larock, VCH, 1989, p. 6ff. The transformation is effected by means of catalytic hydrogenation, for example using a palladium on carbon catalyst and hydrogen or a hydrogen donor, or by the use of diimide or diborane.

Using the above procedures, but employing, in place of the bromophenyl reagent, 57.1, different bromo-substituted aryl or heteroaryl alkoxy hydroxylamines, and/or different dialkyl alkenyl phosphonates, the products analogous to the compounds, 57.4, 57.6 and 57.7 are obtained.

Example 58



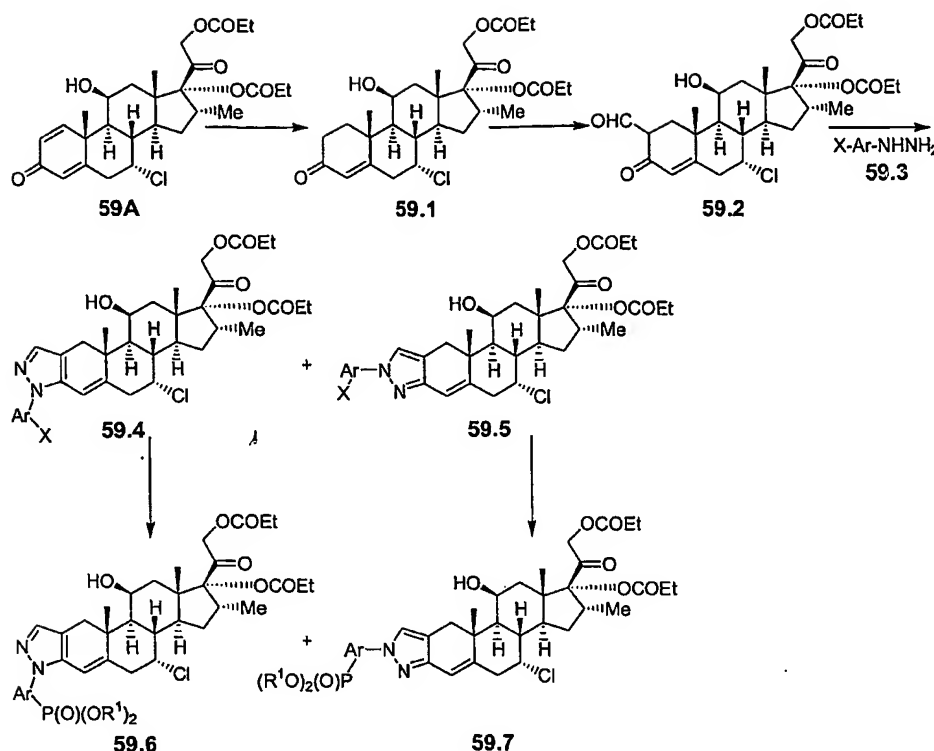
The preparation of phosphonates in which the phosphonate is attached by means of an imino group is illustrated above. In this procedure, the substrate, 55.2, is reacted with a dialkyl 3-aminophenyl phosphonate, 58.1 (*J. Med. Chem.*, 1984, 27, 654), to give, after deprotection and side chain acylation, the imine product, 58.2. The reaction is conducted in a hydrocarbon solvent such as toluene or xylene, at reflux temperature, in the presence of a basic catalyst such as sodium methoxide, or an acid catalyst such as p-toluenesulfonic acid, under azeotropic conditions, to give the product, 58.2.

Using the above procedures, but employing, in place of the 3-aminophenyl phosphonate, 58.1, different amino-substituted aryl or heteroaryl phosphonates, products analogous to 58.2 are obtained.

An alternative method for the preparation of phosphonates, 58.3, where the phosphonate is attached by means of an oximino group starts with the

dienone, **55.2**, is illustrated above. The dienone is reacted with hydroxylamine to yield, after deprotection and side chain acylation, the oxime, **58.3**. The reaction of steroidal 1,4-dien-3-ones with hydroxylamine is described in *J. Steroid Bioch.*, 1976, 7, 795; the reaction is performed between equimolar amounts of the reactants in a polar organic solvent such as pyridine or methanol, optionally in the presence of acetic acid or sodium acetate. The oxime is then reacted with a dialkyl 3-hydroxyphenyl phosphonate, **58.4** (Epsilon), in a Mitsunobu reaction, to yield the substituted oxime, **58.5**. The preparation of aromatic ethers and thioethers by means of the Mitsunobu reaction is described, for example, in "Comprehensive Organic Transformations," by R. C. Larock, VCH, 1989, p. 448, and in "Advanced Organic Chemistry," Part B, by F.A. Carey and R. J. Sundberg, Plenum, 2001, p. 153-4 and in *Org. React.*, 1992, 42, 335. The phenol and the hydroxy or mercapto component are reacted together in an aprotic solvent such as, for example, tetrahydrofuran, in the presence of a dialkyl azodicarboxylate and a triarylphosphine, to afford the ether or thioether products. The procedure is also described in *Org. React.*, 1992, 42, 335-656.

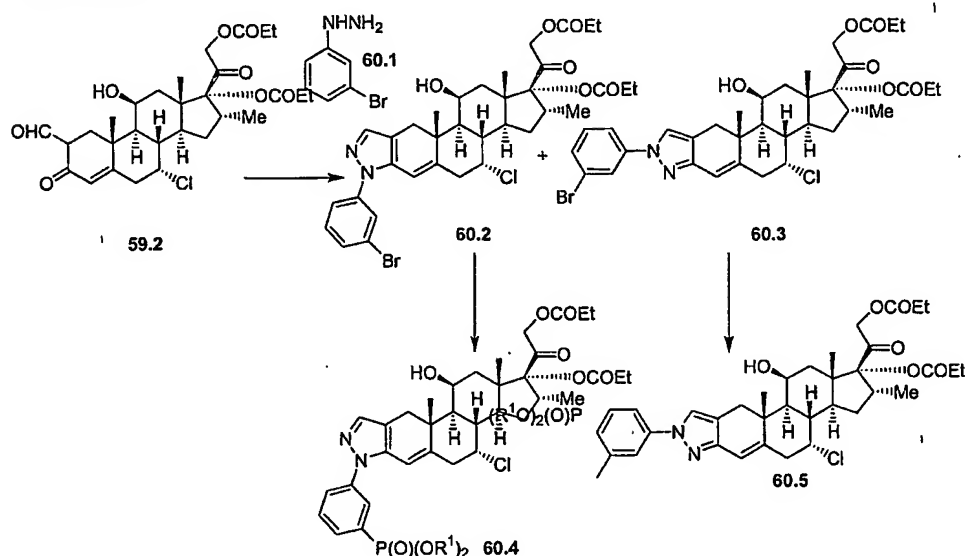
Using the above procedures, but employing, in place of the phosphonate, **58.4**, different dialkyl hydroxy-substituted aryl or heteroaryl phosphonates, the products analogous to **58.5** are obtained.

Example 59

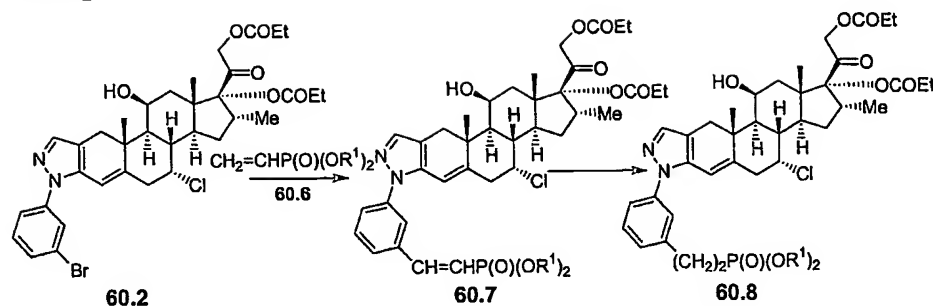
The preparation of the phosphonate esters in which the phosphonate group is attached to the 1' or 2' position of the pyrazole ring, by means of an aromatic or heteroaromatic group, a heteroatom and a variable carbon chain is shown above. In this procedure, Aclometasone dipropionate, **59A**, is reduced to afford the 1,2-dihydro product, **59.1**. The catalytic hydrogenation reaction is effected by the use of tris(triphenylphosphine)rhodium (I) chloride, for example as described in *J. Med. Chem.*, **2001**, 44, 602. The product is then reacted with ethyl formate and a base such as sodium hydride, in an inert solvent such as toluene or dimethylformamide, as described in Australian Patent Application 275950409, to afford the 2-formyl product, **59.2**. Optionally, the substrate, **59A**, is protected, for example, as described above in Example 55, prior to the formylation reaction, as described in *J. Am. Chem. Soc.*, **1964**, 86, 1520. The 2-formyl product is then reacted with an aryl or heteroaryl hydrazine, **59.3**, in which the substituent X is either a phosphonate group or a group which is subsequently transformed into a phosphonate-containing substituent. For example, X is dialkylphosphono, bromo, hydroxy, amino, carboxyl and the like. The reaction yields the isomeric 2'- and 1'-aryl pyrazoles, **59.4** and **59.5**. The

ring-forming reaction is performed between equimolar amounts of the reactants in an acidic solvent such as acetic acid, as described in *J. Am. Chem. Soc.*, 1964, 86, 1520. The pyrazoles, 59.4 and 59.5, are then transformed, for example by the procedures described in Examples 60 - 65, respectively into the phosphonates, 59.6 and 59.7.

Example 60

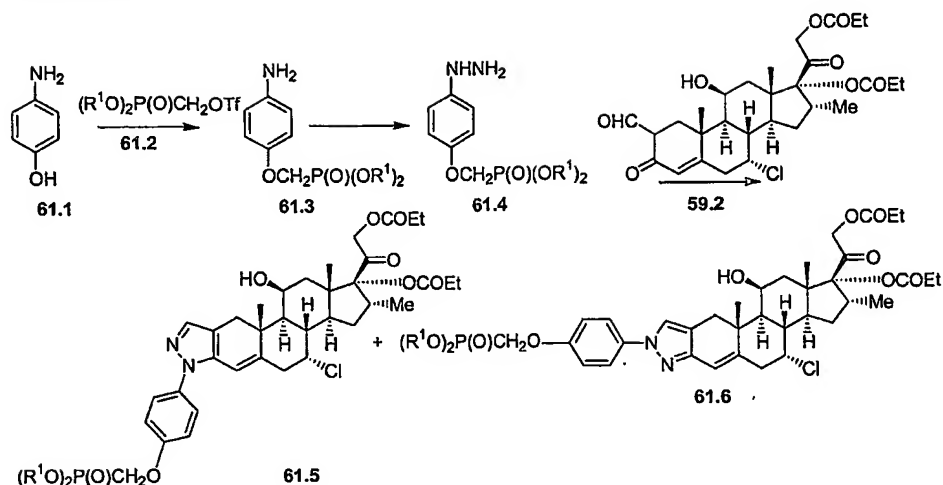


The preparation of phosphonates in which the phosphonate is attached by means of a phenyl group is illustrated above. In this sequence, the ketoaldehyde, 59.2, is reacted, as described above, with 3-bromophenylhydrazine, 60.1 (Fluka), to give the isomeric pyrazole products, 60.2 and 60.3. The products are then reacted, as described above, with a dialkyl phosphite $\text{HP(O)(OR}^1)_2$ and a palladium catalyst, to afford respectively the phosphonates, 60.4 and 60.5. Using the above procedures, but employing, in place of 3-bromophenylhydrazine, different bromoaryl or bromoheteroaryl hydrazines, 59.3, the corresponding products, 59.6 and 59.7 are obtained.

Example 60A

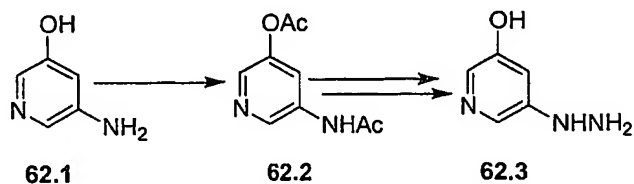
The preparation of phosphonates in which the phosphonate is attached by means of an aromatic or heteroaromatic group and a saturated or unsaturated alkyl chain is illustrated above. In this procedure, the bromophenyl-substituted pyrazole, **60.2**, is coupled in a Heck reaction, as described above, with, for example a dialkyl vinyl phosphonate, **60.6** (Aldrich), to give the unsaturated phosphonate product, **60.7**. Optionally, the product is reduced, as described above, to give the saturated analog, **60.8**. Application of the above procedures to the isomeric bromophenyl pyrazole, **60.3**, affords the products isomeric with, **60.7** and **60.8**.

Using the above procedures, but employing, in place of the phosphonate, **60.6**, different dialkyl alkenyl phosphonates, and/or different bromoaryl or heteroaryl pyrazoles, **59.4** or **59.5** ($X = \text{Br}$), the products analogous to **60.7** and **60.8** are obtained.

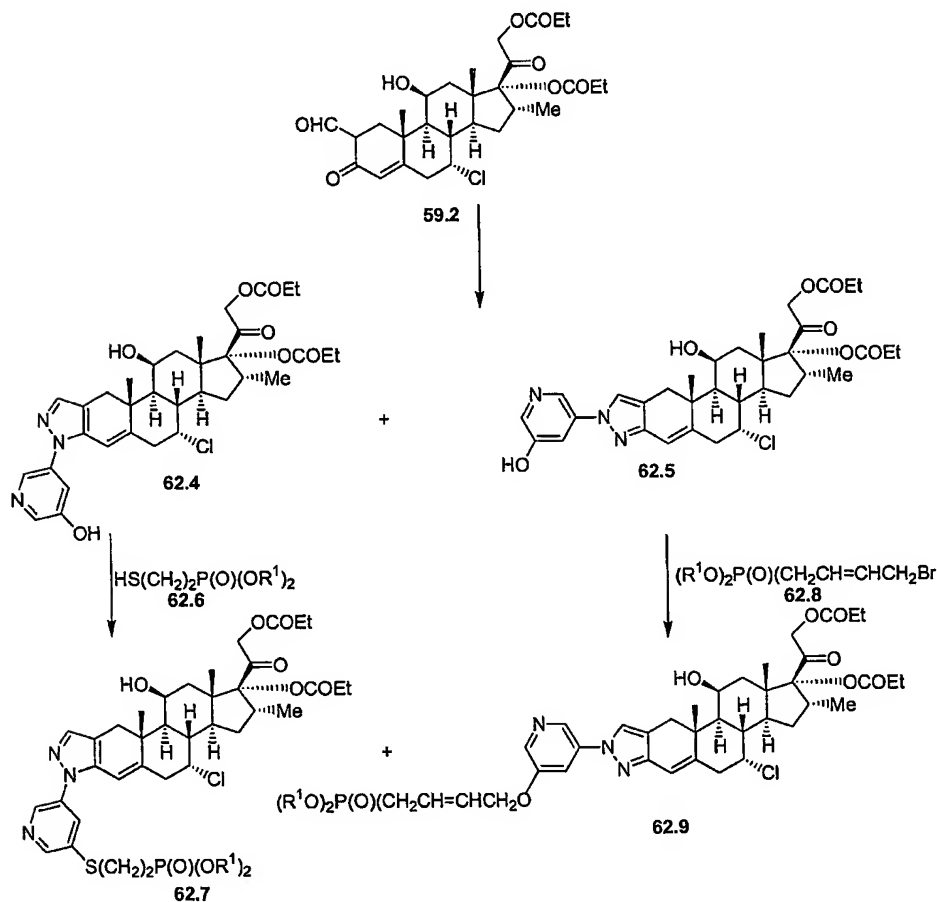
Example 61

The preparation of phosphonates in which the phosphonate is attached by means of an aryl or heteroaryl group and an alkoxy chain is illustrated above. In this procedure, 4-aminophenol, **61.1**, is reacted in dimethylformamide solution at ambient temperature with a dialkyl trifluoromethanesulfonyloxymethyl phosphonate, **61.2** (*Tetrahedron Lett.*, 1986, 27, 1477), and potassium carbonate to give the ether, **61.3**. The product is then converted into the corresponding hydrazine, **61.4**, by means of a diazotization reaction in aqueous ethanolic hydrochloric acid, followed by reduction of the diazonium chloride with tin(II) chloride, as described in *J. Med. Chem.*, 2001, 44, 4031. The hydrazine is then reacted, as described above, with the ketoaldehyde, **59.2**, to form the isomeric pyrazoles, **61.5** and **61.6**.

Using the above procedures, but employing, in place of the triflate, **61.2**, different dialkylphosphono alkyl bromides or triflates, and/or different aromatic or heteroaromatic hydroxyamines, the products analogous to **61.5** and **61.6** are obtained.

Example 62

The preparation of phosphonates in which the phosphonate is attached by means of a pyridyl group, a heteroatom, and a variable carbon chain is shown herein. In this procedure, 3-amino-5-hydroxypyridine, **62.1**, is converted, by reaction with acetic anhydride, into the diacetyl analog, **62.2**. The product is then transformed by diazotization and reduction, as described above, into the hydrazine, **62.3**.



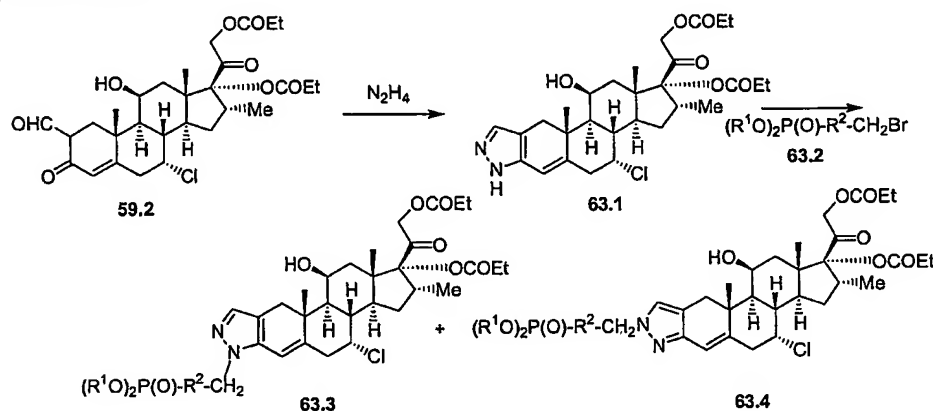
The hydrazine, **62.3**, is then reacted with the ketoaldehyde, **59.2**, to give the isomeric pyrazoles, **62.4** and **62.5**. The 2'-pyridyl product, **62.4**, is reacted in a Mitsunobu reaction, as described above, with a dialkyl mercaptoethyl phosphonate, **62.6** (*Zh. Obshchei. Khim.*, 1973, 43, 2364), to afford the thioether, **62.7**. Application of this procedure to the isomeric phenol, **62.5**, affords the product isomeric to, **62.7**.

Alternatively, the isomeric phenol, **62.5**, is reacted, in dimethylformamide solution at ca. 80°, with one molar equivalent of a dialkyl bromo-

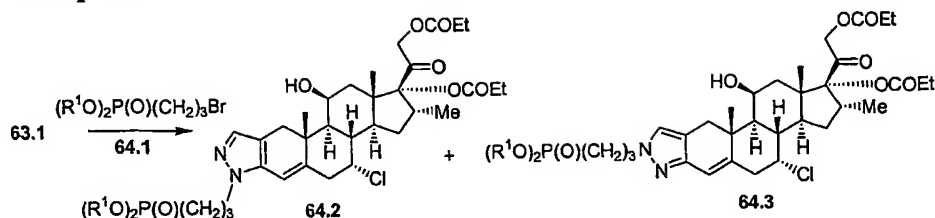
butenyl phosphonate, **62.8** (*J. Med. Chem.*, **1992**, *35*, 1371), and cesium carbonate, to prepare the phosphonate, **62.9**. Application of this procedure to the isomeric phenol, **62.4**, affords the product isomeric with, **62.9**.

Using the above procedures, but employing, in place of the thiol, **62.6**, or the bromide, **62.8**, different thiols, alcohols or bromides, and/or different phenols, **59.4** or **59.5**, in which X is OH, the corresponding products analogous to **62.7** and **62.9** are obtained.

Example 63



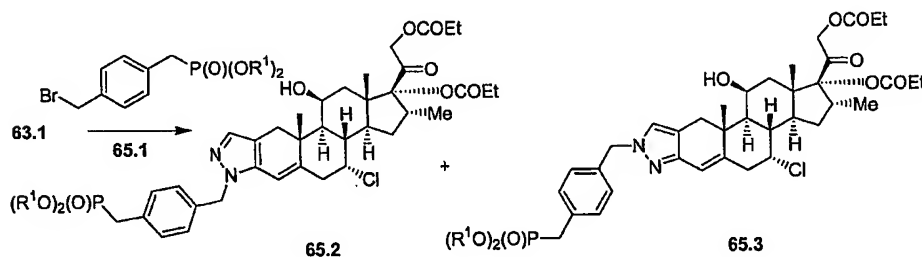
The preparation of the phosphonate esters in which the phosphonate group is attached by means of a variable carbon linkage is illustrated above. In this procedure, the ketoaldehyde, **59.2**, is reacted with hydrazine, to afford the pyrazole derivative, **63.1**. The reaction of steroidal 2-formyl-3-ketones with hydrazine is described in *J. Am. Chem. Soc.*, **1964**, *86*, 1520. The reaction is performed in acetic acid at ambient temperature. The resulting pyrazole is then reacted with a dialkyl bromomethyl phosphonate, **63.2**, in which R^2 is as defined above, to produce the isomeric 2' and 1' alkylation products, **63.3** and **63.4**, respectively. The alkylation of substituted pyrazoles is described, for example, in "Heterocyclic Chemistry," by T. L. Gilchrist, Longman, **1992**, p. 309.

Example 64

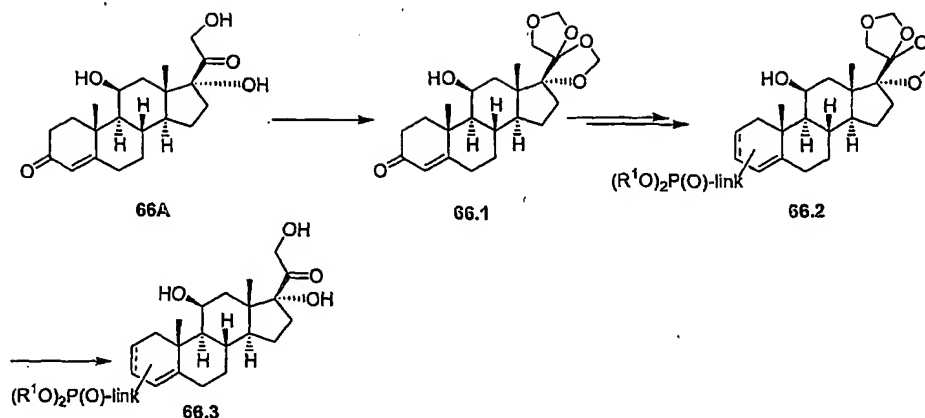
The pyrazole, **63.1**, is reacted, in dimethylformamide solution at ca. 90°, with a dialkyl bromopropyl phosphonate, **64.1** (Aldrich), and a base such as dimethylaminopyridine or lithium hexamethyldisilazide, to yield the isomeric alkylation products, **64.2** and **64.3**.

Example 65

Pyrazole, **63.1**, is reacted, as described above, with a dialkyl 4-bromomethyl benzyl phosphonate, **65.1** (*Tet.*, 1998, 54, 9341), to give the products, **65.2** and **65.3**.

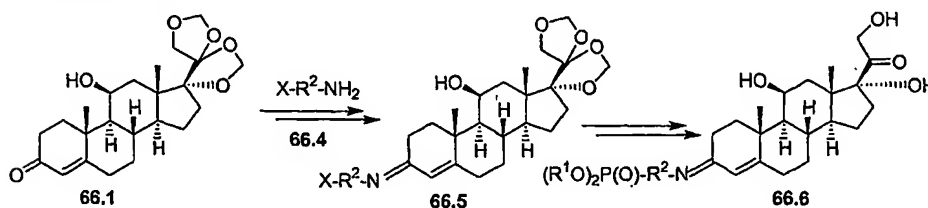
**Example 66**

The structures of Hydrocortisone, **66A** (U.S. Patent No. 2,602,769) and the phosphonate esters are shown below, in which the substituent R¹ is H, alkyl, alkenyl, aryl or aralkyl. These compounds incorporate a phosphonate moiety (R¹O)₂P(O) connected to the nucleus by means of a variable linking group, designated as "link" in the attached structures. A general protection-deprotection sequence in which the steroid side-chain is protected as a bis-methylenedioxy (BMD) moiety is shown below.



Hydrocortisone, **66A**, is reacted with paraformaldehyde and an acid catalyst such as hydrochloric acid, as described "Protective Groups in Organic Synthesis," by T.W. Greene and P.G.M. Wuts, Wiley, Second Edition 1990, p. 223, to yield the BMD derivative, **66.1**. The phosphonate moiety is then introduced, using the procedures described below, to produce the phosphonate ester, **66.2**. The BMD moiety is then hydrolyzed, for example by treatment with 50% aqueous acetic acid, as described in "Protective Groups in Organic Synthesis," by T.W. Greene and P.G.M. Wuts, Wiley, Second Edition 1990, p. 223, to afford the triol, **66.3**.

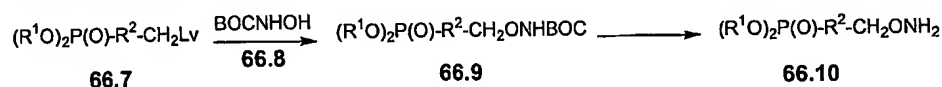
Example 66A



The BMD-protected derivative, **66.1**, is reacted with an amine or hydroxylamine, **66.4**, in which R^2 is an alkyl, alkenyl, cycloalkyl or cycloalkenyl group, optionally incorporating a heteroatom O, S or N, or a functional group such as an amide, ester, oxime, sulfoxide or sulfone etc, or an optionally substituted aryl, heteroaryl or aralkyl group, optionally incorporating a heteroatom O, S or N, and X is either a phosphonate group or a group which is subsequently converted into a phosphonate-containing substituent. For example, X is dialkylphosphono, bromo, hydroxy, amino, carboxy and the like. The

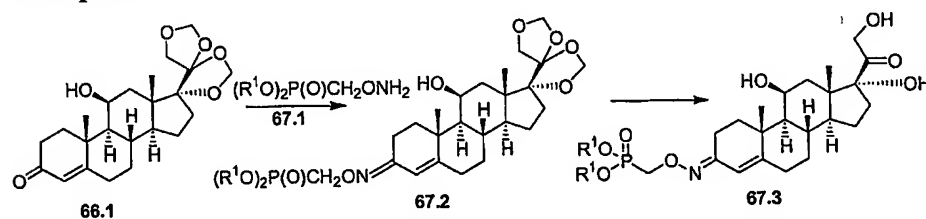
reaction is conducted between equimolar amounts of the reactants in an aprotic solvent such as pyridine or xylene, or in an alcoholic solvent such as ethanol, optionally in the presence of an acid catalyst, to give the imine or oxime, 66.5. The preparation of oximes of steroidal 3-ketones is described in *Anal. Bioch.*, 1978, 86, 133 and in *J. Mass. Spectrom.*, 1995, 30, 497. In cases in which X is not dialkylphosphono, the substituent X is converted, using the methods described below; into a phosphonate-containing substituent; the BMD-protected side-chain is then removed to afford the triol, 66.6.

Example 66B



The preparation of hydroxylamine ethers incorporating a phosphonate group is illustrated above. In this procedure, a phosphonate, 66.7, in which Lv is a leaving group such as bromo or trifluoromethylsulfonyloxy, is reacted with BOC-hydroxylamine, 66.8 (Aldrich), to produce the ether, 66.9. The reaction is conducted between equimolar amounts of the reactants in a polar solvent such as dimethylformamide or tetrahydrofuran, in the presence of a base such as potassium hydroxide or dimethylaminopyridine. Deprotection, for example by treatment with trifluoroacetic acid, then gives the hydroxylamine ether, 66.10.

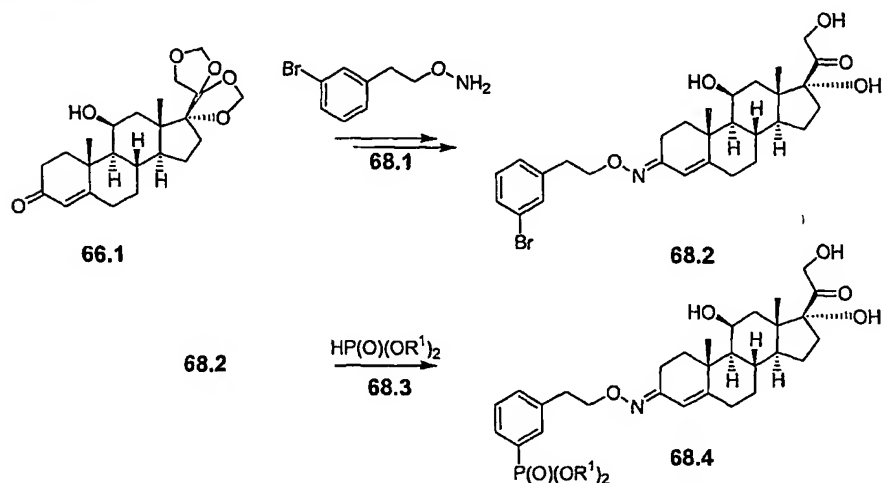
Example 67



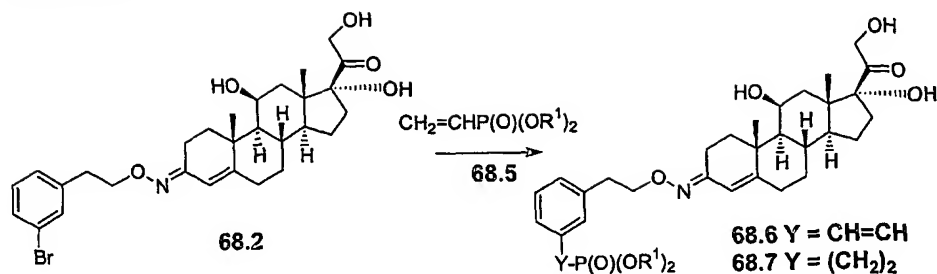
The substrate, 66.1, is reacted with a dialkyl phosphonomethyl hydroxylamine, 67.1, prepared as described above from a dialkyl trifluoromethylsulfonyloxymethyl phosphonate (*Tetrahedron Lett.*, 1986, 27, 1477) and BOC-hydroxylamine, to afford the oxime, 67.2, which is deprotected to afford the triol, 67.3. The oxime forming reaction is performed at ambient temperature in ethanol-acetic acid solution between equimolar amounts of the reactants. Using

the above procedures, but employing, in place of the hydroxylamine ether, 67.1, different oxime ethers, 66.10, the corresponding products, 66.6 are obtained.

Example 68



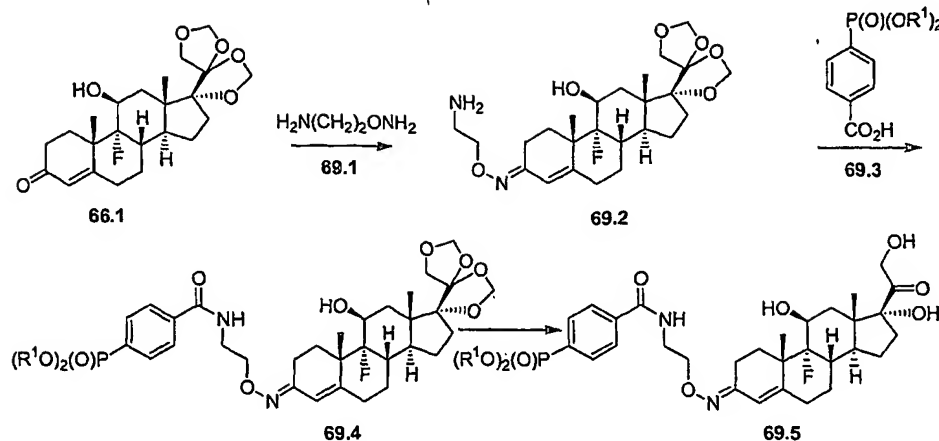
The preparation of compounds in which the phosphonate group is attached by means of a phenyl ethoxy group is illustrated above. In this procedure, the enone, 66.1, is reacted, as described above, with O-(3-bromophenyl)ethyl hydroxylamine, 68.1, prepared as described above from 2-(3-bromophenyl)ethyl bromide (French Patent FR 1,481,052), to give, after deprotection of the side-chain, the oxime, 68.2. The product is then reacted, in the presence of a palladium catalyst, with a dialkyl phosphite, 68.3, to afford the phosphonate, 68.4. The preparation of arylphosphonates by means of a coupling reaction between aryl bromides and dialkyl phosphites is described in *J. Med. Chem.*, 1992, 35, 1371. The reaction is performed in an inert solvent such as toluene, in the presence of a base, such as triethylamine, and a catalytic amount of tetrakis(triphenylphosphine)palladium(0).

Example 68A

Alternatively, the bromo compound, **68.2**, is coupled with a dialkyl vinylphosphonate, **68.5** (Aldrich), to afford the phosphonate, **68.6**. The coupling of aryl halides with olefins by means of the Heck reaction is described, for example, in "Advanced Organic Chemistry," by F. A. Carey and R. J. Sundberg, Plenum, 2001, p. 503ff and in *Acc. Chem. Res.*, 1979, 12, 146.

The aryl bromide and the olefin are coupled in a polar solvent such as dimethylformamide or dioxan, in the presence of a palladium(0) catalyst such as tetrakis(triphenylphosphine)palladium(0) or palladium(II) catalyst such as palladium(II) acetate, and optionally in the presence of a base such as triethylamine or potassium carbonate. Optionally, the styrenoid double bond present in the product **68.6** is reduced, for example by reaction with diimide, to produce the saturated analog, **68.7**. The reduction of olefinic bonds is described in "Comprehensive Organic Transformations," by R. C. Larock, VCH, 1989, p. 6ff. The transformation is effected by means of catalytic hydrogenation, for example using a palladium on carbon catalyst and hydrogen or a hydrogen donor, or by the use of diimide or diborane.

Using the above procedures, but employing, in place of the bromophenyl ethoxy reagent, **68.1**, different bromo-substituted aryl or heteroaryl alkoxy hydroxylamines, and/or different dialkyl alkenyl phosphonates, the products analogous to the compounds, **68.4**, **68.6** and **68.7** are obtained.

Example 69

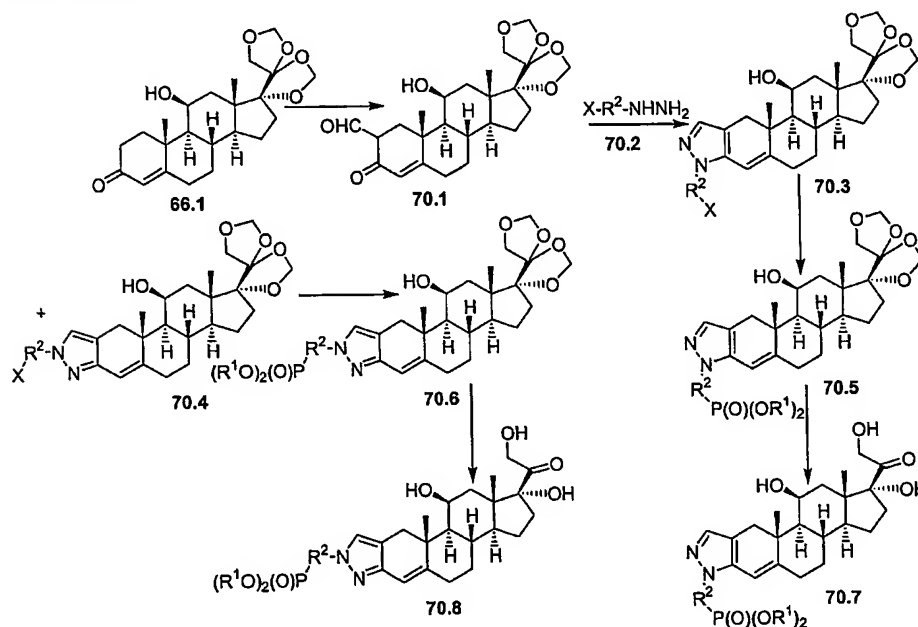
The enone, **66.1**, is reacted with O-(2-aminoethyl)hydroxylamine, **69.1**, (*Pol. J. Chem.*, **1981**, *55*, 1163) to yield the oxime, **69.2**. The reaction of steroidal 1,4-dien-3-ones with substituted hydroxylamines is described in *J. Steroid Bioch.*, **1976**, *7*, 795. The reaction is performed between equimolar amounts of the reactants in a polar organic solvent such as pyridine or methanol, optionally in the presence of acetic acid or sodium acetate. The oxime is then coupled with a dialkyl 4-carboxyphenyl phosphonate, **69.3** (Epsilon), to yield the amide oxime, **69.4**. The preparation of amides from carboxylic acids and derivatives is described, for example, in *Organic Functional Group Preparations*, by S.R.Sandler and W. Karo, Academic Press, 1968, p. 274, and *Comprehensive Organic Transformations*, by R. C. Larock, VCH, 1989, p. 972ff. The carboxylic acid is reacted with the amine in the presence of an activating agent, such as, for example, dicyclohexylcarbodiimide or diisopropylcarbodiimide, optionally in the presence of, for example, hydroxybenztriazole, N-hydroxysuccinimide or N-hydroxypyridone, in a non-protic solvent such as, for example, pyridine, DMF or dichloromethane, to afford the amide.

Alternatively, the carboxylic acid (*e.g.*, dialkyl 4-carboxyphenyl phosphonate, **69.3**) is first converted into an activated derivative such as the acid chloride, anhydride, mixed anhydride, imidazolide and the like, and then reacted with the amine, in the presence of an organic base such as, for example, pyridine, to afford the amide. The conversion of a carboxylic acid into the corresponding acid chloride is effected by treatment of the carboxylic acid with a

reagent such as, for example, thionyl chloride or oxalyl chloride in an inert organic solvent such as dichloromethane, optionally in the presence of a catalytic amount of dimethylformamide. The amide product, 69.4, is then converted, as described herein, into the triol, 69.5.

Using the above procedures, but employing, in place of the hydroxylamine, 69.1, different amino-substituted hydroxylamines, and/or different carboxy-substituted phosphonates, the products analogous to 69.5 are obtained.

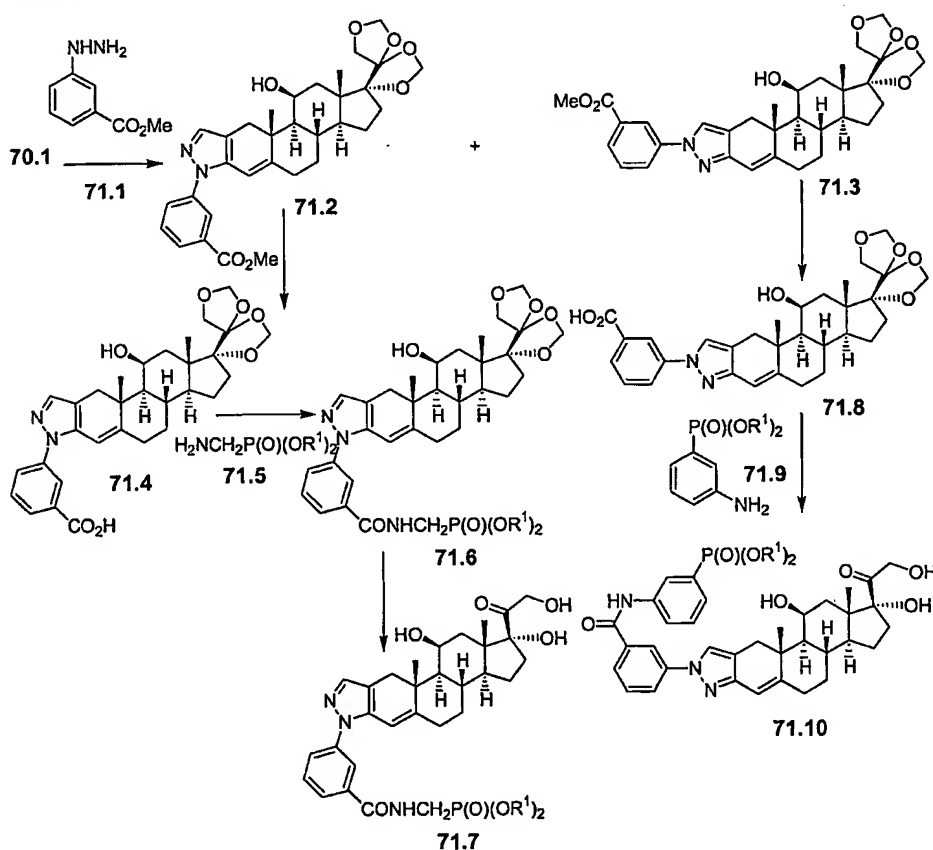
Example 70



The preparation of the phosphonate esters in which the phosphonate group is attached to the 1' or 2' position of the pyrazole ring, by means of a variable carbon chain is illustrated above. In this procedure, the BMD-protected enone, 66.1, is reacted with ethyl formate and a base such as sodium hydride, in an inert solvent such as toluene or dimethylformamide, as described in *J. Am. Chem. Soc.*, 1964, 86, 1520, to afford the 2-formyl product, 70.1. This compound is then reacted with an alkyl, aralkyl, aryl or heteroaryl hydrazine, 70.2, wherein R² is alkyl, aralkyl, aryl, or heteroaryl, and in which the substituent X is either a phosphonate group or a group which is subsequently transformed into a phosphonate-containing substituent. For example, X is

dialkylphosphono, bromo, hydroxy, amino, carboxyl and the like. The reaction yields the isomeric 2'- and 1'-aryl pyrazoles, **70.3** and **70.4**. The pyrazole-forming reaction is performed between equimolar amounts of the reactants in an acidic solvent such as acetic acid, as described in *J. Am. Chem. Soc.*, 1964, 86, 1520. The pyrazoles, **70.3** and **70.4**, are then transformed, for example, using procedures described herein, via the BMD-protected intermediates, **70.5** and **70.6**, into the phosphonates, **70.7** and **70.8**.

Example 71

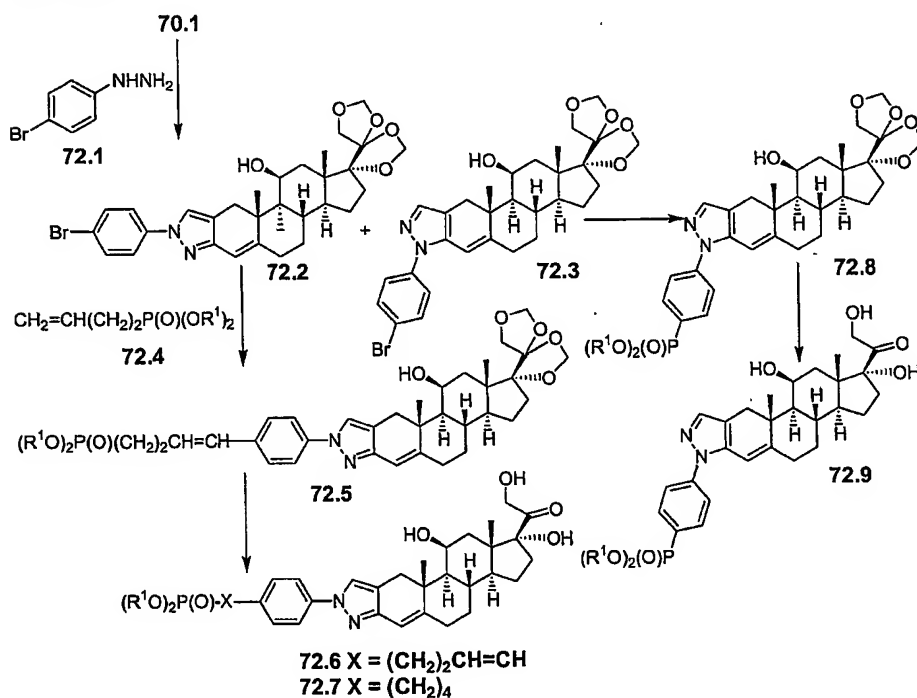


The preparation of phosphonates in which the phosphonate moiety is attached by means of a phenyl ring and an amide linkage is illustrated above. In this procedure, the ketoaldehyde, **70.1**, is reacted, with 3-carbomethoxyphenylhydrazine, **71.1** (Apin), to give the pyrazoles, **71.2** and **71.3**. The 2'-substituted isomer, **71.2**, is then reacted with one molar equivalent of lithium hydroxide in aqueous dimethoxyethane, to produce the carboxylic acid, **71.4**. The acid is then

coupled, as described above, with a dialkyl aminomethyl phosphonate, **71.5**, (Interchim) to give the amide, **71.6**, deprotection then affords the triol, **71.7**. Alternatively, the 1'-substituted pyrazole, **71.3**, is hydrolyzed, as described above, to the carboxylic acid, **71.8**. The product is then coupled with a dialkyl 3-aminophenyl phosphonate, **71.9** (*J. Med. Chem.*, 1984, 27, 654), to yield after deprotection the triol amide, **71.10**.

Using the above procedures, but employing, in place of the carbomethoxyphenyl hydrazine, **71.1**, different carbomethoxy-substituted aralkyl, aryl or heteroaryl alkoxy hydrazines, and/or different dialkyl amino-substituted phosphonates, the products analogous to the compounds, **71.7** and **71.10** are obtained.

Example 72



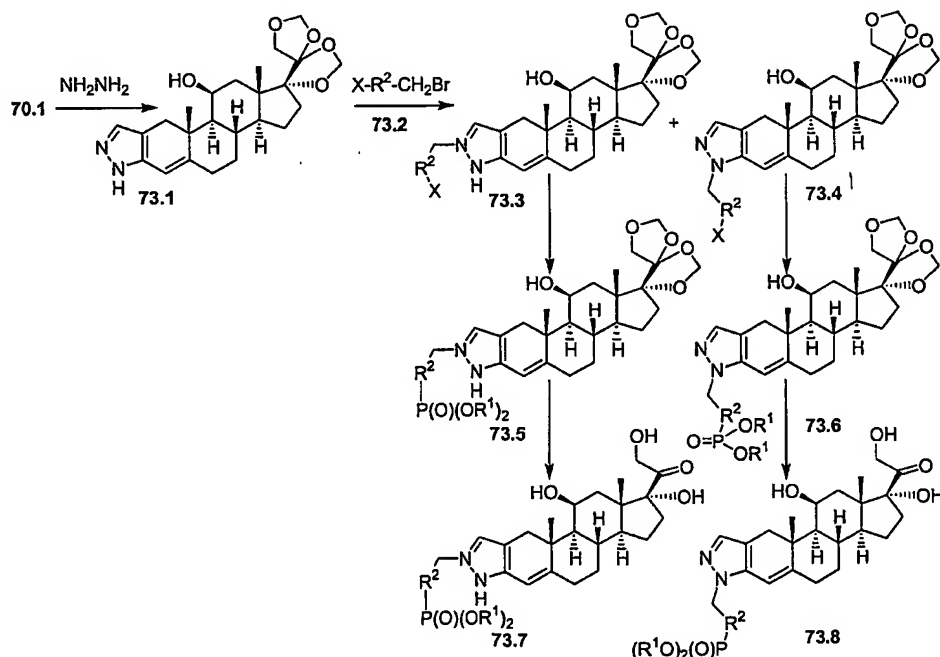
The preparation of the phosphonate esters in which the phosphonate group is attached by means of a variable carbon linkage is illustrated above. In this procedure, the ketoaldehyde, **70.1**, is reacted, as described above, with 4-bromophenyl hydrazine, **72.1** (*J. Organomet. Chem.*, 1999, 62, 581), to produce the pyrazoles, **72.2** and **72.3**. The 1'-substituted isomer, **72.2**, is coupled, as described herein, in the presence of a palladium catalyst, with a dialkyl butenyl

phosphonate, **72.4** (*Org. Lett.*, **2001**, *3*, 217), to give the phosphonate, **72.5**. The product is then deprotected to afford the triol, **72.6**. Optionally, the styrenoid double bond present in the product, **72.6**, is reduced, as described above, to produce the saturated analog, **72.7**.

Alternatively, the 2'-substituted pyrazole, **72.3**, is coupled, in the presence of a palladium catalyst, with a dialkyl phosphite to prepare the phosphonate, **72.8**, which is deprotected to give the triol, **72.9**. The preparation of arylphosphonates by means of a coupling reaction between aryl bromides and dialkyl phosphites is described in *J. Med. Chem.*, **1992**, *35*, 1371. This reaction is performed in an inert solvent such as toluene, in the presence of a base such as triethylamine and tetrakis(triphenylphosphine)-palladium(0).

Using the above procedures, but employing, in place of the bromophenyl hydrazine, **72.1**, different bromo-substituted aralkyl, aryl or heteroaryl alkoxy hydrazines, and/or different dialkyl alkenyl phosphonates, the products analogous to the compounds, **72.6**, **72.7** and **72.9** are obtained.

Example 73

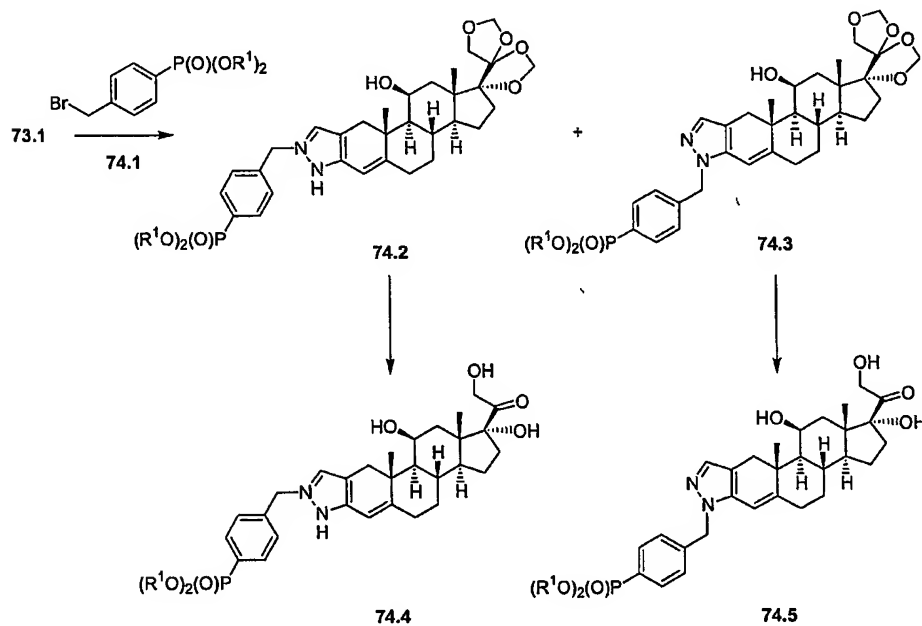


The preparation of the phosphonate esters in which the phosphonate group is attached by means of a variable carbon linkage is illustrated above. In this procedure, the ketoaldehyde, **70.1**, is reacted with hydrazine, to afford the

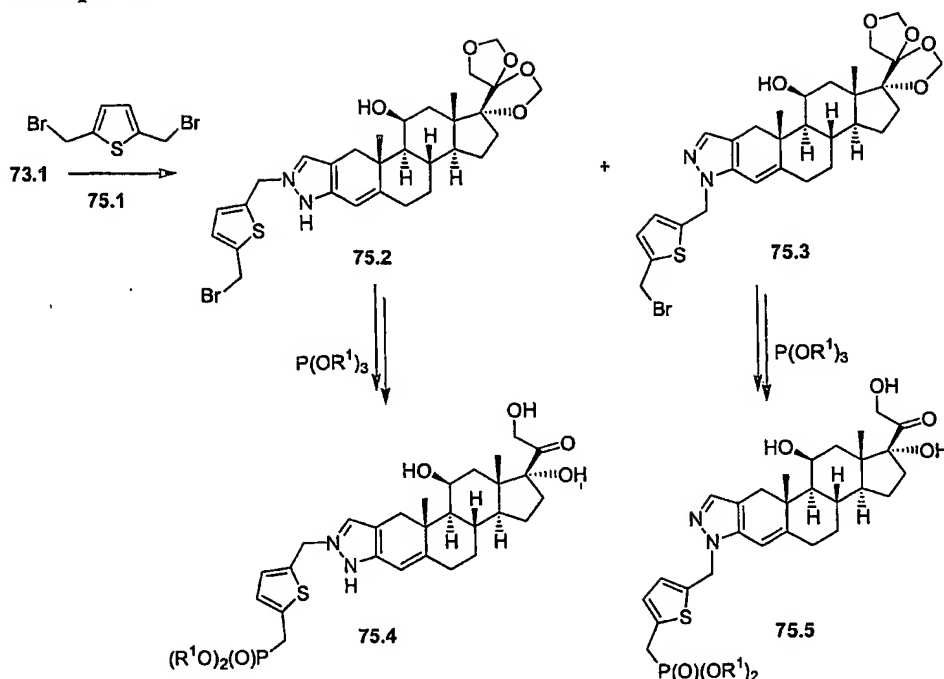
pyrazole derivative, **73.1**. The reaction of steroidal 2-formyl-3-ketones with hydrazine is described in *J. Am. Chem. Soc.*, **1964**, 86, 1520. The reaction is performed in ethanol at reflux temperature. The pyrazole product is then reacted with a bromomethyl compound, **73.2**, in which R^2 and X are as defined above, to yield the alkylation products, **73.3** and **73.4**. The alkylation of substituted pyrazoles is described, for example, in "Heterocyclic Chemistry," by T. L. Gilchrist, Longman, **1992**, p. 309.

The reaction is performed between equimolar amounts of the substrates in a polar solvent such as dimethylformamide or tetrahydrofuran, in the presence of a base such as dimethylaminopyridine, lithium hexamethyldisilazide and the like. The products, **73.3** and **73.4**, are converted into the phosphonates, **73.5** and **73.6**, except in cases where X is dialkylphosphono, using the procedures described herein. Deprotection affords the triols, **73.7** and **73.8**.

Example 74



The pyrazole, **73.1**, is reacted, as described above, with one molar equivalent of a dialkyl 4-(bromomethyl)phenyl phosphonate, **74.1** (WO 2003/042150), to give the alkylated pyrazoles, **74.2** and **74.3**. Deprotection then yields the triols, **74.4** and **74.5**.

Example 75

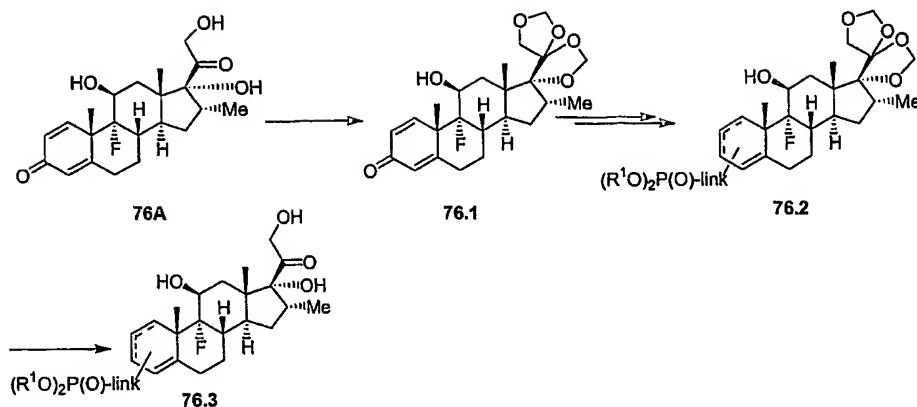
The pyrazole, 73.1, is reacted, as described above, with 2,5-bis(bromomethyl)thiophene, 75.1 (*Tet.*, 1999, 55, 4709), to give the pyrazoles, 75.2 and 75.3. The products are subjected to an Arbuzov reaction, in which the bromomethyl substituent is converted into the dialkyl phosphonomethyl substituent, by reaction with a trialkyl phosphite at 120°, to prepare, after deprotection of the side chain, the phosphonates, 75.4 and 75.5. The Arbuzov reaction is described in *Handb. Organophosphorus Chem.*, 1992, 115-72. In the procedure, the substrate is heated at from 60° to about 160° with a five to fifty-fold molar excess of the trialkyl phosphite.

Using the above procedures, but employing, in place of the dibromide, 75.1, different dibromides, the products analogous to 75.4 and 75.5 are obtained.

Example 76

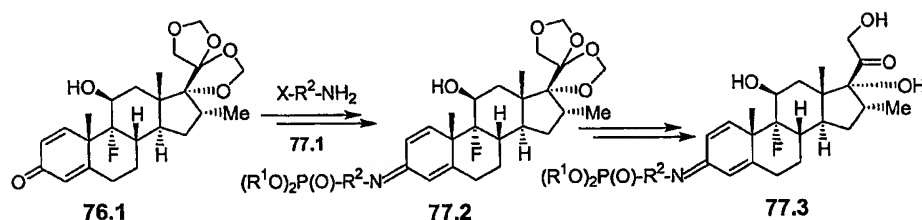
Dexamethasone, 76A, (U.S. Patent No. 3,007,923) analogs where the substituent R^1 is H, alkyl, alkenyl, aryl or aralkyl are prepared by the procedures below. The compounds 76.1 – 76.3 incorporate a phosphonate moiety $(R^1O)_2P(O)$ connected to the nucleus by means of a variable linking group, designated as "link" in the attached structures.

Scheme M1



Dexamethasone, **76A**, is reacted with paraformaldehyde and an acid catalyst such as hydrochloric acid, as described in "Protective Groups in Organic Synthesis," by T.W. Greene and P.G.M. Wuts, Wiley, Second Edition 1990, p. 223, to yield the BMD derivative, **76.1**. The phosphonate moiety is then introduced, using the procedures described below, to produce the phosphonate ester, **76.2**. The BMD moiety is then hydrolyzed, for example by treatment with 50% aqueous acetic acid, as described in "Protective Groups in Organic Synthesis," by T.W. Greene and P.G.M. Wuts, Wiley, Second Edition 1990, p. 223, to afford the triol, **76.3**.

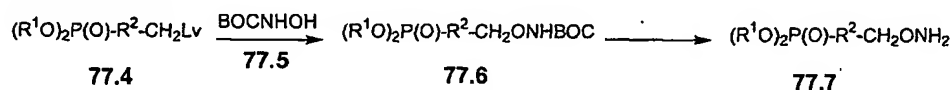
Example 77



The preparation of phosphonates in which the phosphonate is attached by means of an imino or iminoxy group and a variable carbon chain is illustrated above. In this procedure, the BMD-protected derivative, **76.1**, is reacted with an amine or hydroxylamine, **77.1**, in which R^2 is an alkyl, alkenyl, cycloalkyl or cycloalkenyl group, optionally incorporating a heteroatom O, S or N, or a functional group such as an amide, ester, oxime, sulfoxide or sulfone etc, or an optionally substituted aryl, heteroaryl or aralkyl group, optionally incorporating

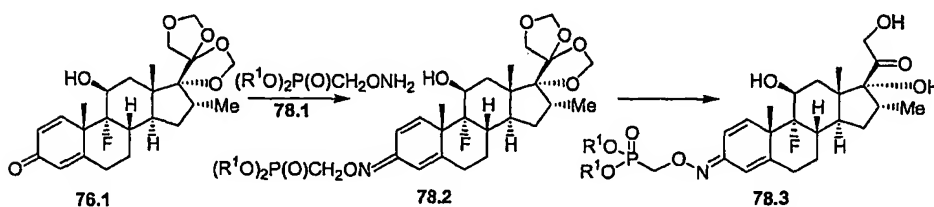
a heteroatom O, S or N, and X is either a phosphonate group or a group which is subsequently converted into a phosphonate-containing substituent. For example, X is dialkylphosphono, bromo, hydroxy, amino, carboxy and the like. The reaction is conducted between equimolar amounts of the reactants in an aprotic solvent such as pyridine or xylene, or in an alcoholic solvent such as ethanol, optionally in the presence of an acid catalyst, to give the imine or oxime. The preparation of oximes of steroidal 3-ketones is described in *Anal. Bioch.*, 1978, 86, 133 and in *J. Mass. Spectrom.*, 1995, 30, 497. The BMD-protected side-chain compound, 77.2, is then converted, as described in Scheme M1 in Example 76, into the triol, 77.3.

Example 77A



The preparation of hydroxylamine ethers incorporating a phosphonate group is illustrated above. In this procedure, a phosphonate, 77.4, in which Lv is a leaving group such as bromo or trifluoromethylsulfonyloxy, is reacted with BOC-hydroxylamine, 77.5 (Aldrich), to produce the ether, 77.6. The reaction is conducted between equimolar amounts of the reactants in a polar solvent such as dimethylformamide or tetrahydrofuran, in the presence of a base such as potassium hydroxide or dimethylaminopyridine. Deprotection, for example by treatment with trifluoroacetic acid, then gives the hydroxylamine ether, 77.7.

Example 78

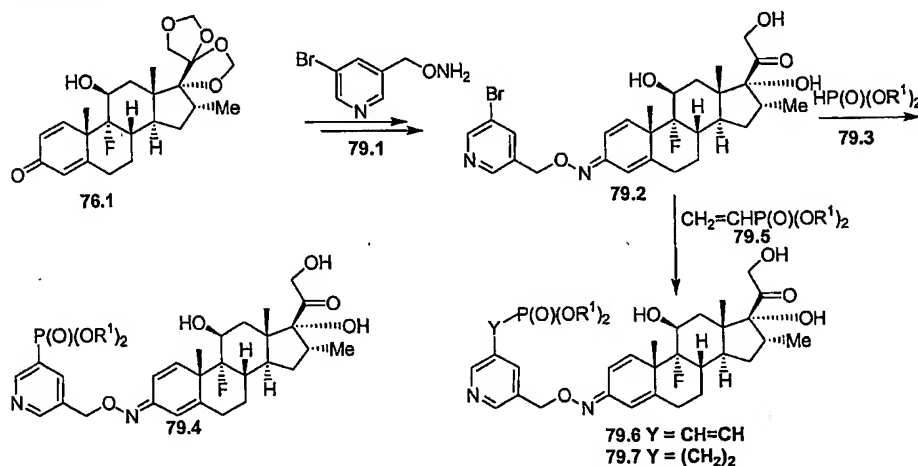


The preparation of phosphonates in which the phosphonate is attached by means of an iminoxy group. In this procedure, the substrate, 76.1, is reacted with a dialkyl phosphonomethyl hydroxylamine, 78.1, prepared as described above from a dialkyl trifluoromethylsulfonyloxymethyl phosphonate (*Tetrahedron Lett.*, 1986, 27, 1477) and BOC-hydroxylamine, to afford the

oxime, **78.2**, which is deprotected to afford the triol, **78.3**. The oxime forming reaction is performed at ambient temperature in ethanol-acetic acid solution between equimolar amounts of the reactants.

Using the above procedures, but employing, in place of the hydroxylamine ether, **78.1**, different oxime ethers, **77.1**, the corresponding products, **77.3** are obtained.

Example 79



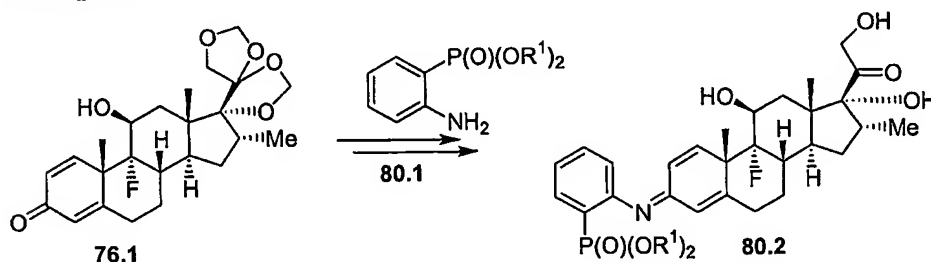
The preparation of compounds in which the phosphonate group is attached by means of a pyridyl methoxy group is illustrated above. In this procedure, the dienone, **76.1**, is reacted, as described above, with O-(3-bromo-5-pyridylmethyl)hydroxylamine, **79.1**, prepared as described above from 3-bromo-5-bromomethylpyridine (WO 95/28400), to give, after deprotection of the side-chain, the oxime, **79.2**. The product is then reacted, in the presence of a palladium catalyst, with a dialkyl phosphite, **79.3**, to afford the phosphonate, **79.4**. The preparation of arylphosphonates by means of a coupling reaction between aryl bromides and dialkyl phosphites is described in *J. Med. Chem.*, **1992**, 35, 1371. The reaction is performed in an inert solvent such as toluene, in the presence of a base such as triethylamine and a catalytic amount of tetrakis-(triphenylphosphine)palladium(0).

Alternatively, the bromo compound, **79.2**, is coupled with a dialkyl vinylphosphonate, **79.5** (Aldrich), to afford the phosphonate, **79.6**. The coupling of aryl halides with olefins by means of the Heck reaction is described, for example, in "Advanced Organic Chemistry," by F. A. Carey and R. J. Sundberg,

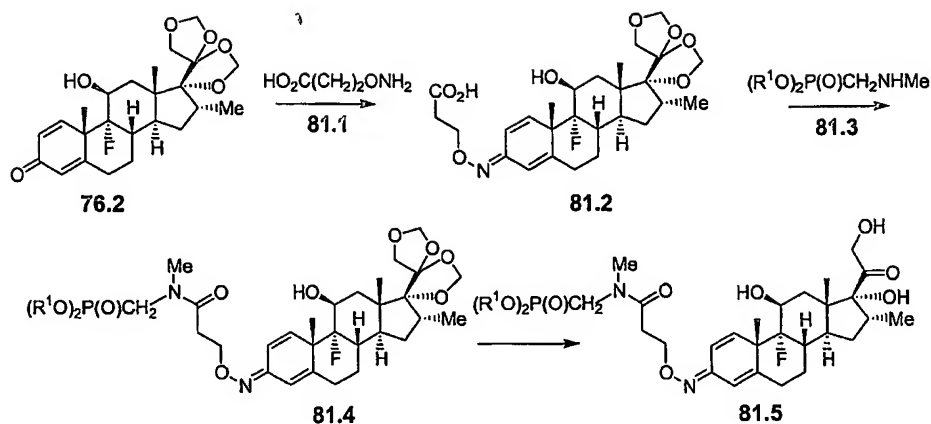
Plenum, 2001, p. 503ff and in *Acc. Chem. Res.*, 1979, 12, 146. The aryl bromide and the olefin are coupled in a polar solvent such as dimethylformamide or dioxan, in the presence of a palladium(0) catalyst such as tetrakis(triphenylphosphine)palladium(0) or palladium(II) catalyst such as palladium(II) acetate, and optionally in the presence of a base such as triethylamine or potassium carbonate. Optionally, the styrenoid double bond present in the product, 79.6, is reduced, for example by reaction with diimide, to produce the saturated analog, 79.7. The reduction of olefinic bonds is described in "Comprehensive Organic Transformations," by R. C. Larock, VCH, 1989, p. 6ff. The transformation is effected by means of catalytic hydrogenation, for example using a palladium on carbon catalyst and hydrogen or a hydrogen donor, or by the use of diimide or diborane.

Using the above procedures, but employing, in place of the bromopyridyloxy reagent, 79.1, different bromo-substituted aryl or heteroaryl alkoxy hydroxylamines, and/or different dialkyl alkenyl phosphonates, the products analogous to the compounds, 79.4, 79.6 and 79.7 are obtained.

Example 80



The preparation of phosphonates in which the phosphonate is attached by means of an imino group is illustrated above. In this procedure, the substrate, 76.1, is reacted with a dialkyl 2-aminophenyl phosphonate, 80.1, (*Syn.*, 1999, 1368) to give, after deprotection, the imine product, 80.2. The reaction is conducted in a hydrocarbon solvent such as toluene or xylene, at reflux temperature, in the presence of a basic catalyst such as sodium methoxide, or an acid catalyst such as p-toluenesulfonic acid, under azeotropic conditions. Using the above procedures, but employing, in place of the 2-aminophenyl phosphonate, 80.1, different amino-substituted aryl or heteroaryl phosphonates, products analogous to 80.2 are obtained.

Example 81

The preparation of phosphonates in which the phosphonate is attached by means of an oximino group and an amide linkage is illustrated above. In this procedure, the dienone, 76.2, is reacted with O-(2-carboxyethyl)hydroxylamine, 81.1, (*J. Med. Chem.*, 1990, 33, 1423) to yield the oxime, 81.2. The reaction of steroidal 1,4-dien-3-ones with substituted hydroxylamines is described in *J. Steroid Bioch.*, 1976, 7, 795; the reaction is performed between equimolar amounts of the reactants in a polar organic solvent such as pyridine or methanol, optionally in the presence of acetic acid or sodium acetate. The oxime is then reacted with a dialkyl methylaminomethyl phosphonate, 81.3, to yield the amide oxime, 81.4. The preparation of amides from carboxylic acids and derivatives is described, for example, in "Organic Functional Group Preparations," by S.R. Sandler and W. Karo, Academic Press, 1968, p. 274, and "Comprehensive Organic Transformations," by R. C. Larock, VCH, 1989, p. 972ff. The carboxylic acid is reacted with the amine in the presence of an activating agent, such as, for example, dicyclohexylcarbodiimide or diisopropylcarbodiimide, optionally in the presence of, for example, hydroxybenzotriazole, N-hydroxysuccinimide or N-hydroxypyridone, in a non-protic solvent such as, for example, pyridine, DMF or dichloromethane, to afford the amide.

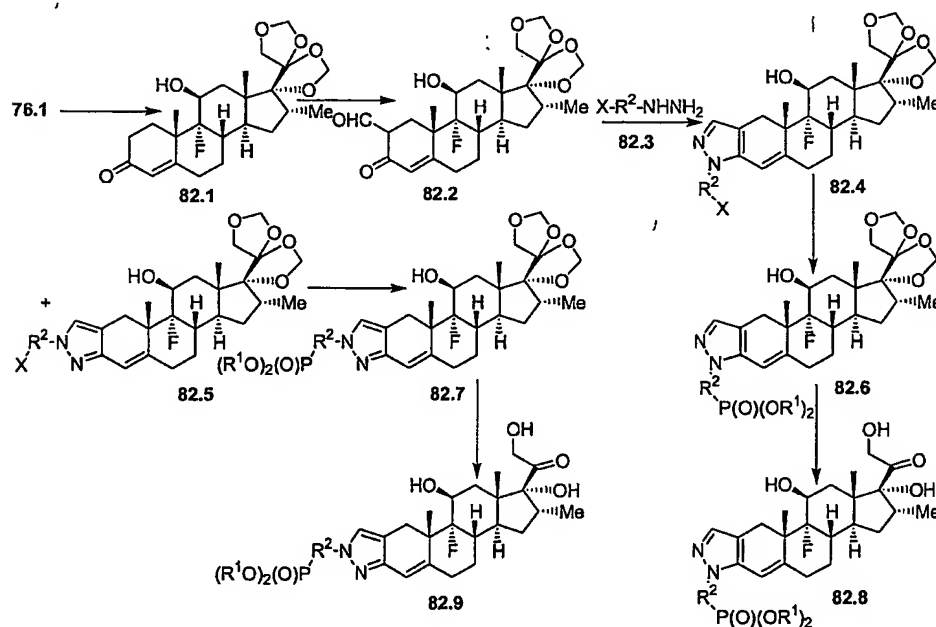
Alternatively, the carboxylic acid is first converted into an activated derivative such as the acid chloride, anhydride, mixed anhydride, imidazolid and the like, and then reacted with the amine, in the presence of an organic base

such as, for example, pyridine, to afford the amide. The conversion of a carboxylic acid into the corresponding acid chloride is effected by treatment of the carboxylic acid with a reagent such as, for example, thionyl chloride or oxalyl chloride in an inert organic solvent such as dichloromethane, optionally in the presence of a catalytic amount of dimethylformamide.

The amide product, **81.4**, is then converted, as described herein, into the triol, **81.5**.

Using the above procedures, but employing, in place of the hydroxylamine, **81.3**, different carboxy-substituted hydroxylamines, and/or different amino-substituted phosphonates, the products analogous to **81.5** are obtained.

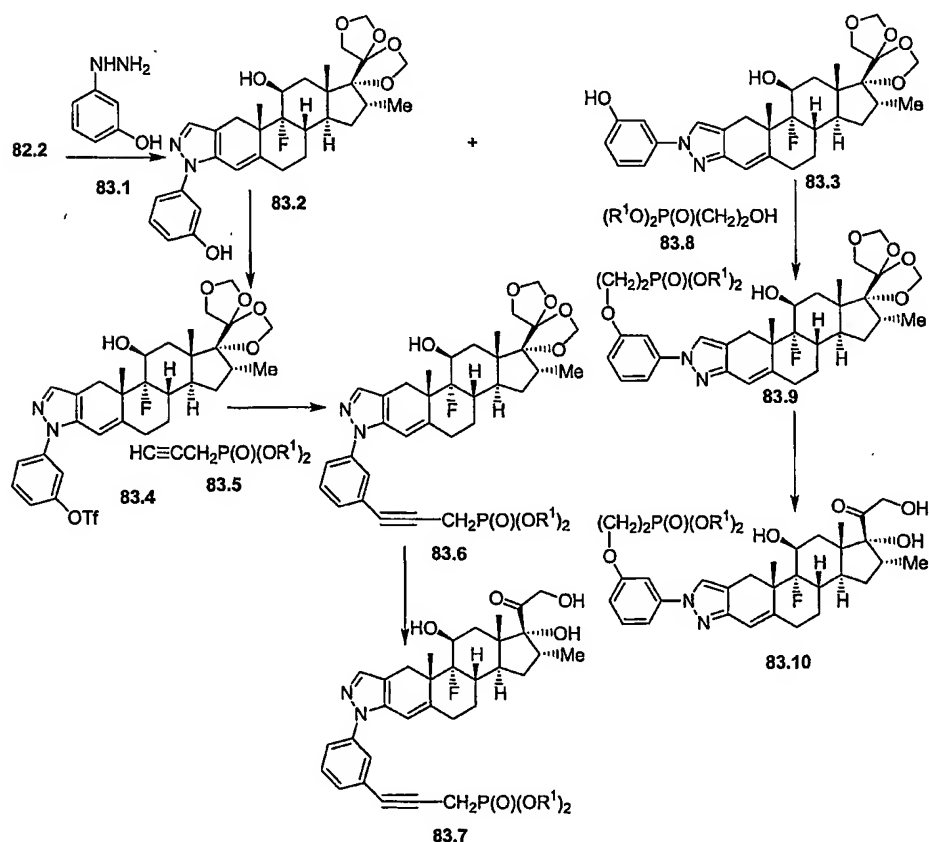
Example 82



The preparation of the phosphonate esters in which the phosphonate group is attached to the 1' or 2' position of the pyrazole ring, by means of an aromatic or heteroaromatic group, a heteroatom and a variable carbon chain is illustrated above. In this procedure, the BMD-protected dienone, **76.1**, is reduced to afford the 1,2-dihydro product, **82.1**. The catalytic hydrogenation reaction is effected by the use of tris(triphenylphosphine)rhodium (I) chloride, for example as described in *J. Med. Chem.*, **2001**, 44, 602. The product is then

reacted with ethyl formate and a base such as sodium hydride, in an inert solvent such as toluene or dimethylformamide, as described in *J. Am. Chem. Soc.*, 1964, 86, 1520, to afford the 2-formyl product, **82.2**. This compound is then reacted with an alkyl, aralkyl, aryl or heteroaryl hydrazine, **82.3**, wherein R^2 is alkyl, aralkyl, aryl or heteroaryl and in which the substituent X is either a phosphonate group or a group which is subsequently transformed into a phosphonate-containing substituent. For example, X is dialkylphosphono, bromo, hydroxy, amino, carboxyl and the like. The reaction yields the isomeric 2'- and 1'-aryl pyrazoles, **82.4** and **82.5**. The pyrazole-forming reaction is performed between equimolar amounts of the reactants in an acidic solvent such as acetic acid, as described in *J. Am. Chem. Soc.*, 1964, 86, 1520. The pyrazoles, **82.4** and **82.5**, are then transformed, for example by the procedures described herein, via the BMD-protected intermediates, **82.6** and **82.7**, into the phosphonates, **82.8** and **82.9**.

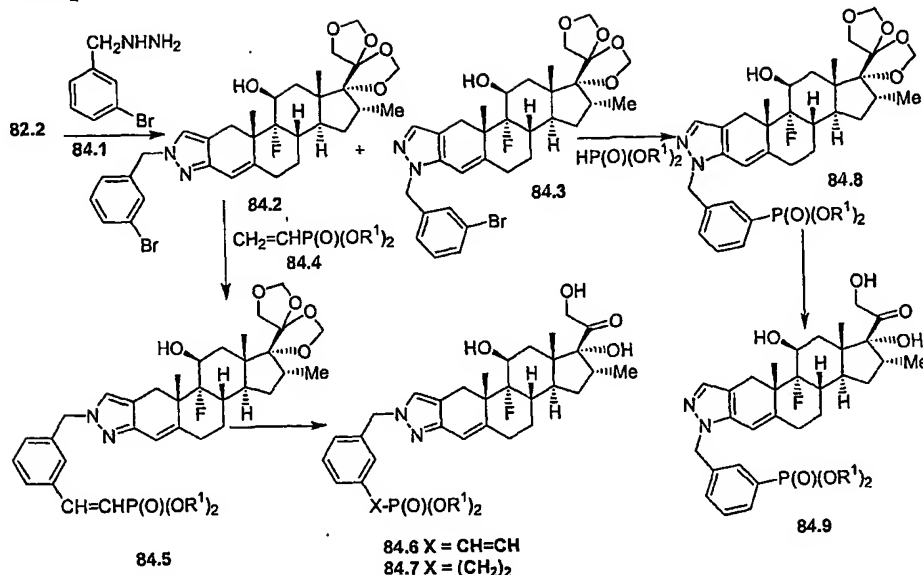
Example 83



The preparation of phosphonates in which the phosphonate is attached by means of a phenyl ring and an alkoxy or an acetylenic linkage is illustrated above. In this procedure, the ketoaldehyde, **82.2**, is reacted, as described above, with 3-hydroxyphenylhydrazine, **83.1** (Japanese Patent No. JP 03011081), to give the pyrazoles, **83.2** and **83.3**. The 2'-substituted isomer, **83.2**, is then reacted in dichloromethane solution at ambient temperature with one molar equivalent of trifluoromethylsulfonyl chloride and dimethylaminopyridine, to yield the triflate, **83.4**. The product is then reacted in toluene solution with a dialkyl propynyl phosphonate, **83.5** (*Syn.*, 1999, 2027), triethylamine and a catalytic amount of tetrakis(triphenylphosphine)palladium (0), to give the acetylenic product, **83.6**. The palladium-catalyzed coupling reaction of aryl triflates with terminal acetylenes is described in WO 02/30930. The BMD protecting group is then removed to yield the triol, **83.7**.

Alternatively, the 1'-substituted pyrazole, **83.3**, is reacted, in a Mitsunobu reaction, with a dialkyl 2-hydroxyethyl phosphonate, **83.8** (Epsilon), to afford the ether, **83.9**. The preparation of aromatic ethers by means of the Mitsunobu reaction is described, for example, in "Comprehensive Organic Transformations," by R. C. Larock, VCH, 1989, p. 448, and in "Advanced Organic Chemistry," Part B, by F.A. Carey and R. J. Sundberg, Plenum, 2001, p. 153-4 and in *Org. React.*, 1992, 42, 335. The phenol and the alcohol component are reacted together in an aprotic solvent such as, for example, tetrahydrofuran, in the presence of a dialkyl azodicarboxylate and a triarylphosphine, to afford the ether or thioether products. The procedure is also described in *Org. React.*, 1992, 42, 335-656. The product, **83.9**, is then deprotected to give the triol, **83.10**.

Using the above procedures, but employing different acetylenic or hydroxyl-substituted phosphonates, the products analogous to **83.7** and **83.10** are obtained. The functionalization procedures are interchangeable between the pyrazole substrates, **83.2** and **83.3**.

Example 84

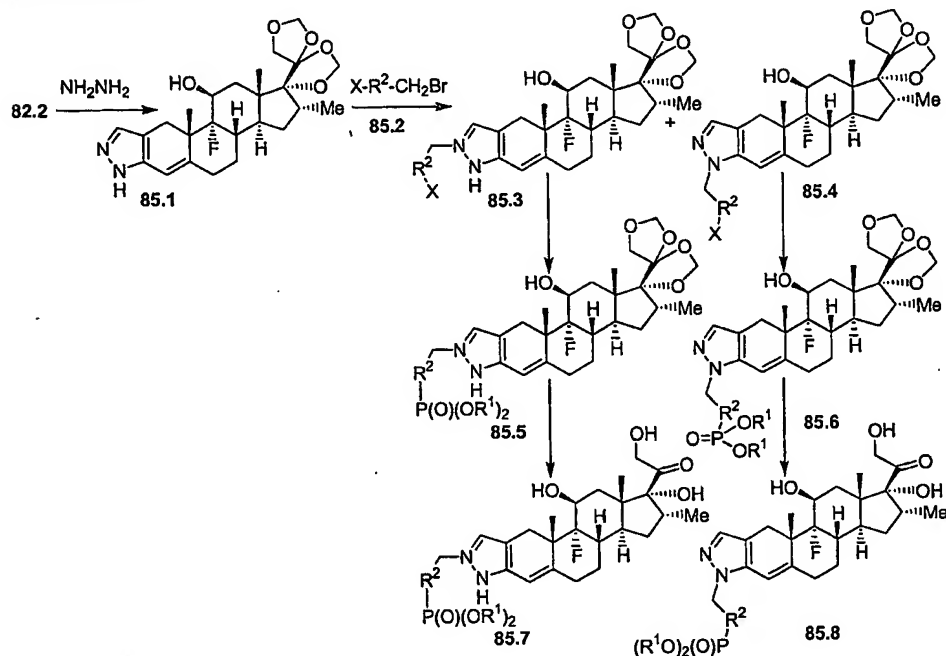
The preparation of the phosphonates in which the phosphonate group is attached by means of a benzyl group and a saturated or unsaturated carbon chain is illustrated above. In this procedure, the ketoaldehyde, **82.2**, is reacted, as described above, with 3-bromobenzyl hydrazine, **84.1** (U.S. Patent No. 4,370,339), to produce the pyrazoles, **84.2** and **84.3**. The 1'-substituted isomer **84.2** is coupled, in the presence of a palladium catalyst, with a dialkyl vinylphosphonate, **84.4** (Aldrich), to give the phosphonate, **84.5**. The product is then deprotected to afford the triol, **84.6**. Optionally, the styrenoid double bond present in the product, **84.6**, is reduced, as described above, to produce the saturated analog, **84.7**.

Alternatively, the 2'-substituted pyrazole, **84.3**, is coupled, in the presence of a palladium catalyst, with a dialkyl phosphite to prepare the phosphonate, **84.8**, which is deprotected to give the triol, **84.9**. The preparation of arylphosphonates by means of a coupling reaction between aryl bromides and dialkyl phosphites is described in *J. Med. Chem.*, **1992**, *35*, 1371. This reaction is performed in an inert solvent such as toluene, in the presence of a base such as triethylamine and tetrakis(triphenylphosphine)palladium(0).

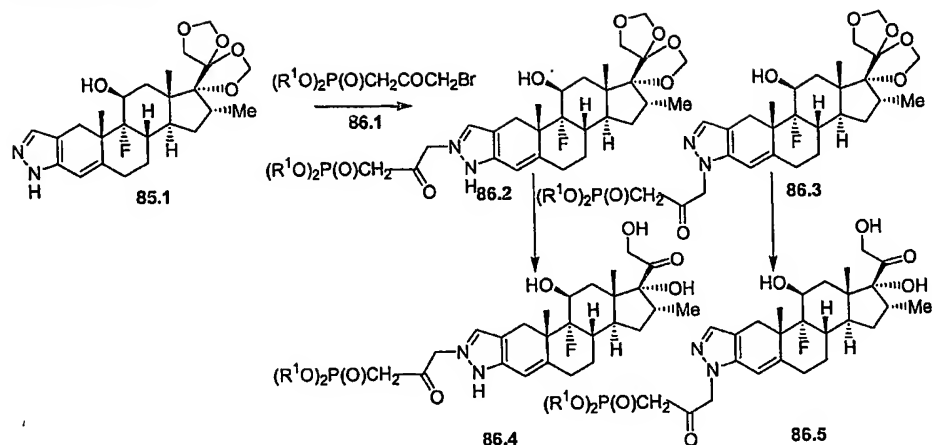
Using the above procedures, but employing, in place of the bromobenzyl reagent, **84.1**, different bromo-substituted aralkyl, aryl or heteroaryl alkoxy

hydrazines, and/or different dialkyl alkenyl phosphonates, the products analogous to the compounds, **84.6**, **84.7** and **84.9** are obtained.

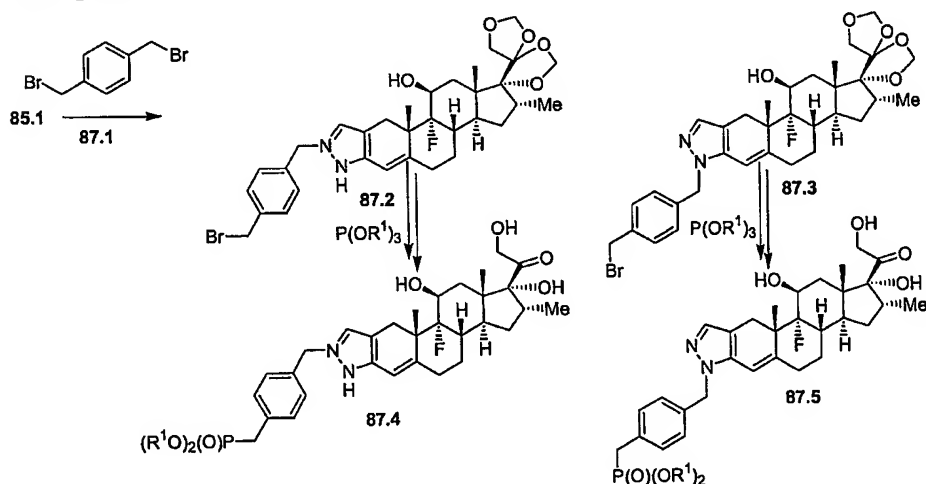
Example 85



The preparation of the phosphonate esters, **85.7** and **85.8**, in which the phosphonate group is attached by means of a variable carbon linkage is illustrated above. In this procedure, the ketoaldehyde, **82.2**, is reacted with hydrazine, to afford the pyrazole derivative, **85.1**. The reaction of steroidal 2-formyl-3-ketones with hydrazine is described in *J. Am. Chem. Soc.*, **1964**, **86**, 1520. The reaction is performed in acetic acid at ambient temperature. The pyrazole product is then reacted with a bromomethyl compound, **85.2**, in which R^2 and X are as defined above, to yield the alkylation products, **85.3** and **85.4**. The alkylation of substituted pyrazoles is described, for example, in "Heterocyclic Chemistry," by T. L. Gilchrist, Longman, **1992**, p. 309. The reaction is performed between equimolar amounts of the substrates in a polar solvent such as dimethylformamide or tetrahydrofuran, in the presence of a base such as dimethylaminopyridine, lithium hexamethyldisilazide and the like. The products, **85.3** and **85.4**, are, except in cases where X is dialkylphosphono, converted into the phosphonates, **85.5** and **85.6**, using the procedures described herein, and deprotection then affords the triols, **85.7** and **85.8**.

Example 86

The pyrazole, 85.1, is reacted, as described above, with one molar equivalent of a dialkyl bromoacetyl phosphonate, 86.1 (*Tet.*, 1978, 34, 649), to give the alkylated pyrazoles, 86.2 and 86.3. Deprotection then yields the triols, 86.4 and 86.5.

Example 87

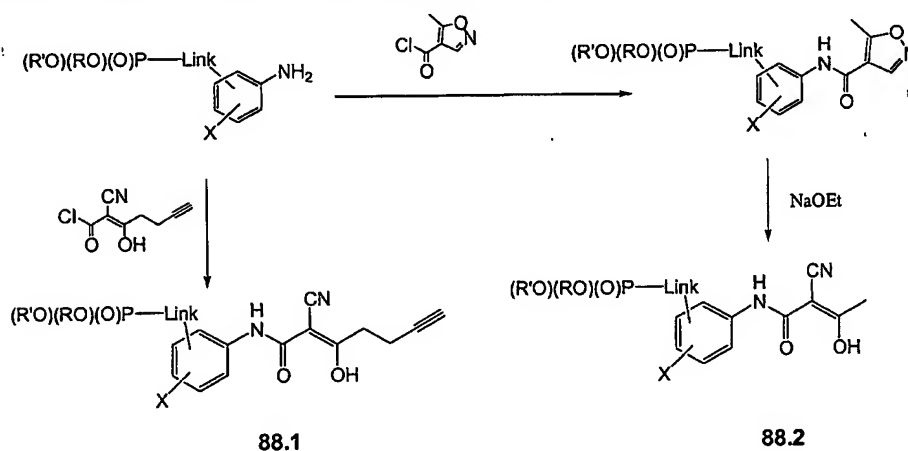
The pyrazole, 85.1, is reacted, as described above, with 1,4-bis(bromomethyl)benzene, 87.1, to give the pyrazoles, 87.2 and 87.3. The products are subjected to an Arbuzov reaction, in which the bromomethyl substituent is converted into the dialkyl phosphonomethyl substituent, by reaction with a trialkyl phosphite at 120°, to prepare, after deprotection of the side chain, the phosphonates, 87.4 and 87.5. The Arbuzov reaction is described in *Handb.*

Organophosphorus Chem., 1992, 115-72.' In the procedure, the substrate is heated at from 60° to about 160° with a five to fifty-fold molar excess of the trialkyl phosphite.

Using the above procedures, but employing, in place of the dibromide, 87.1, different dibromides, the products analogous to 87.4 and 87.5 are obtained.

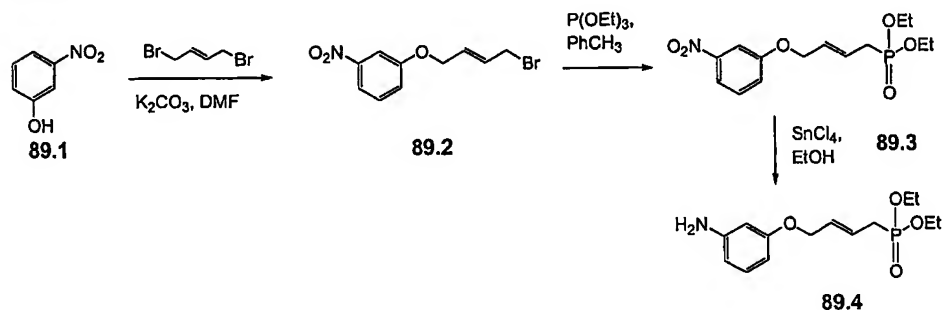
Example 88

A synthetic methodology towards the preparation of phosphonate compounds of Formulae 88.1 and 88.2 is described by Westwood et al, *J. Med. Chem.*, 1996, 39, 4608-4621, according to the general routes outlined below.



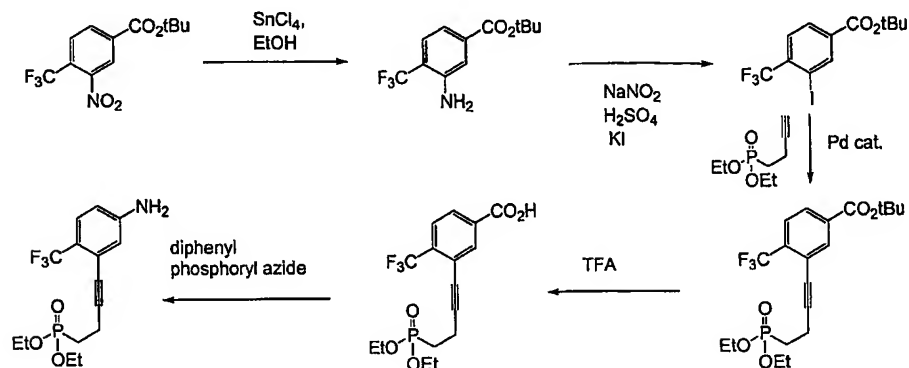
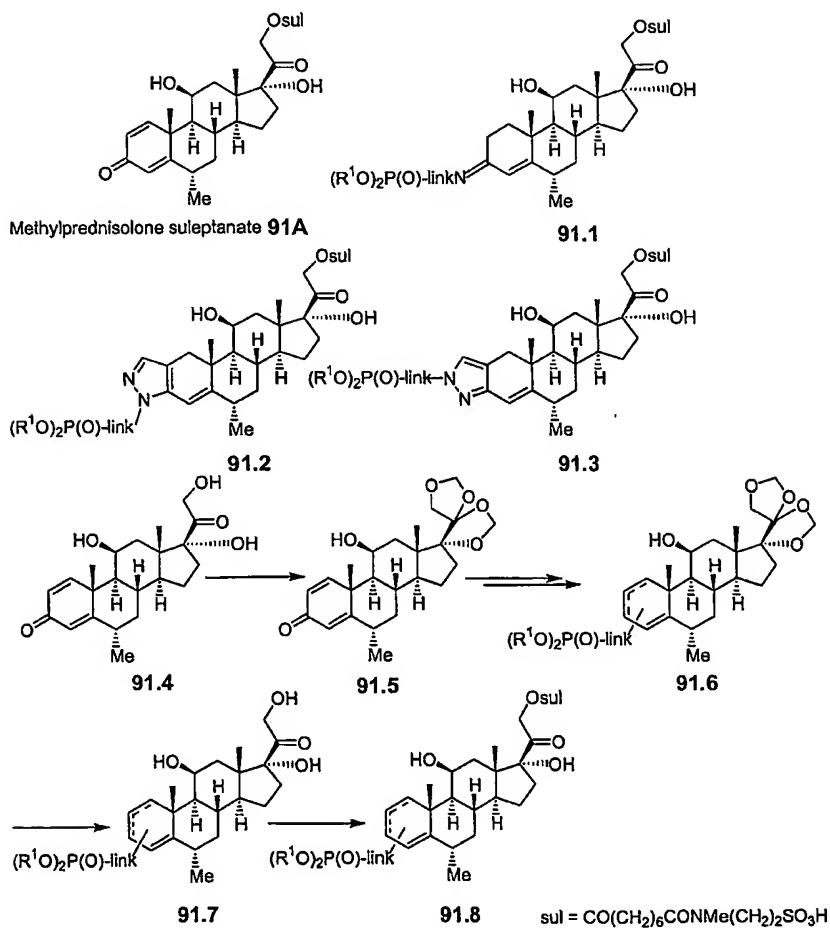
Example 89

The preparation of compounds of the invention having phosphonate groups and intermediate compounds useful for their synthesis are illustrated below.



Example 90

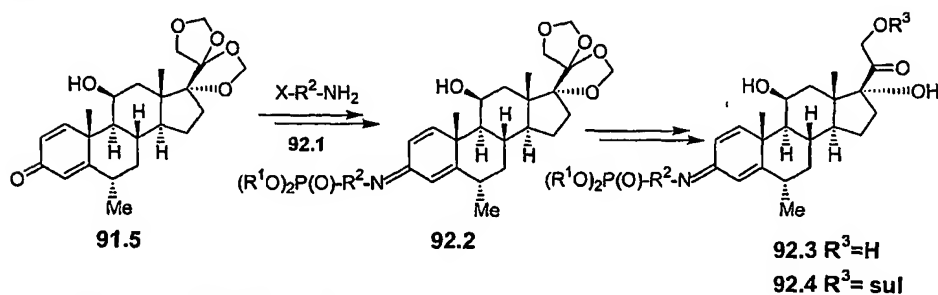
The preparation of compounds of the invention having phosphonate groups and intermediate compounds useful for their synthesis are illustrated below.

**Example 91**

The structures of Methylprednisolone suleptanate, **91A** (WO 89/00558), and the phosphonate esters, **91.1 – 91.3**, are shown above in Example **91**, in which the substituent R^1 is H, alkyl, alkenyl, aryl or aralkyl. The compounds, **91.1 – 91.3**, incorporate a phosphonate moiety $(R^1O)_2P(O)$ connected to the nucleus by means of a variable linking group, designated as "link" in the attached structures. The syntheses of the phosphonate compounds of this invention, **91.1 – 91.3**, and of the intermediate compounds necessary for their synthesis are set forth below.

Methylprednisolone, **91.4**, is reacted with paraformaldehyde and an acid catalyst such as hydrochloric acid, as described in "Protective Groups in Organic Synthesis," by T.W. Greene and P.G.M. Wuts, Wiley, Second Edition 1990, p. 223, to yield the BMD derivative, **91.5**. The phosphonate moiety is then introduced, using the procedures described herein, to produce the phosphonate ester, **91.6**. The BMD moiety is then hydrolyzed, for example by treatment with 50% aqueous acetic acid, as described in "Protective Groups in Organic Synthesis," by T.W. Greene and P.G.M. Wuts, Wiley, Second Edition 1990, p. 223, to afford the triol, **91.7**. The triol is then converted into the 21-suleptanate ester as described in WO 89/00558. In this procedure, a mixed anhydride prepared by reacting suleptanic acid with pivaloyl chloride, in the presence of a base such as triethylamine, is reacted with the 21-hydroxy steroid, **91.7**, to prepare the 21-suleptanate ester, **91.8**.

Example 92

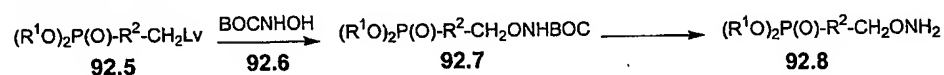


$sul = CO(CH_2)_6CONMe(CH_2)_2SO_3H$

The preparation of phosphonates in which the phosphonate is attached by means of an imino or iminoxy group and a variable carbon chain is illustrated above. In this procedure, the BMD-protected derivative, **91.5**, is reacted with an

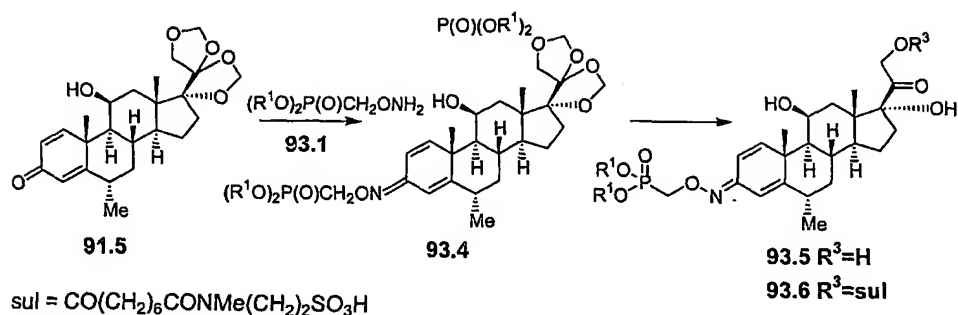
amine or hydroxylamine, **92.1**, in which R^2 is an alkyl, alkenyl, cycloalkyl or cycloalkenyl group, optionally incorporating a heteroatom O, S or N, or a functional group such as an amide, ester, oxime, sulfoxide or sulfone etc, or an optionally substituted aryl, heteroaryl or aralkyl group, optionally incorporating a heteroatom O, S or N, and X is either a phosphonate group or a group which is subsequently converted into a phosphonate-containing substituent. For example, X is dialkylphosphono, bromo, hydroxy, amino, carboxy and the like. The reaction is conducted between equimolar amounts of the reactants in an aprotic solvent such as pyridine or xylene, or in an alcoholic solvent such as ethanol, optionally in the presence of an acid catalyst, to give the imine or oxime. The preparation of oximes of steroidal 3-ketones is described in *Anal. Bioch.*, **1978**, 86, 133 and in *J. Mass. Spectrom.*, **1995**, 30, 497. The BMD-protected side-chain compound, **92.2**, is then converted into the triol, **92.3**, and then to the suleptanate, **92.4**, as described above in Example 91.

Example 92A



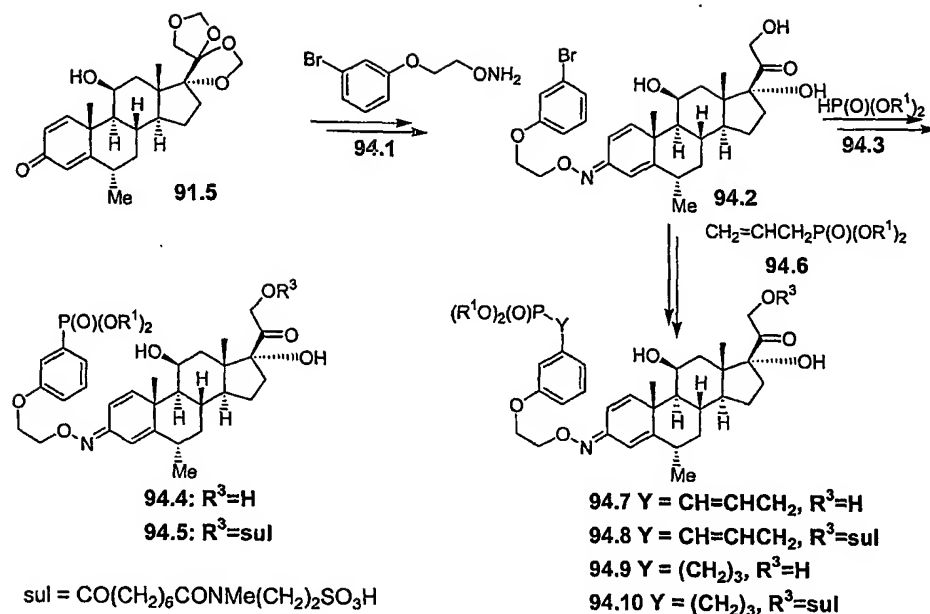
The preparation of hydroxylamine ethers incorporating a phosphonate group is illustrated above. In this procedure, a phosphonate, **92.5**, in which Lv is a leaving group such as bromo or trifluoromethylsulfonyloxy. The phosphonate is reacted with BOC-hydroxylamine, **92.6** (Aldrich), to produce the ether, **92.7**. The reaction is conducted between equimolar amounts of the reactants in a polar solvent such as dimethylformamide or tetrahydrofuran, in the presence of a base such as potassium hydroxide or dimethylaminopyridine. Deprotection, for example by treatment with trifluoroacetic acid, then gives the hydroxylamine ether, **92.8**.

Example 93



The preparation of phosphonates in which the phosphonate is attached by means of an iminoxy group is illustrated above. In this procedure, the substrate, **91.5**, is reacted with a dialkyl phosphonomethyl hydroxylamine, **93.1**, prepared as described above from a dialkyl trifluoromethylsulfonyloxymethyl phosphonate (*Tetrahedron Lett.*, 1986, 27, 1477) and BOC-hydroxylamine, to afford the oxime, **93.4**, which is deprotected to afford the triol, **93.5**, from which the sulleptanate ester, **93.6**, is prepared. The oxime forming reaction is performed at ambient temperature in ethanol-acetic acid solution between equimolar amounts of the reactants.

Using the above procedures, but employing, in place of the hydroxylamine ether, **93.1**, different oxime ethers, **92.1**, the corresponding products, **92.4** are obtained.

Example 94

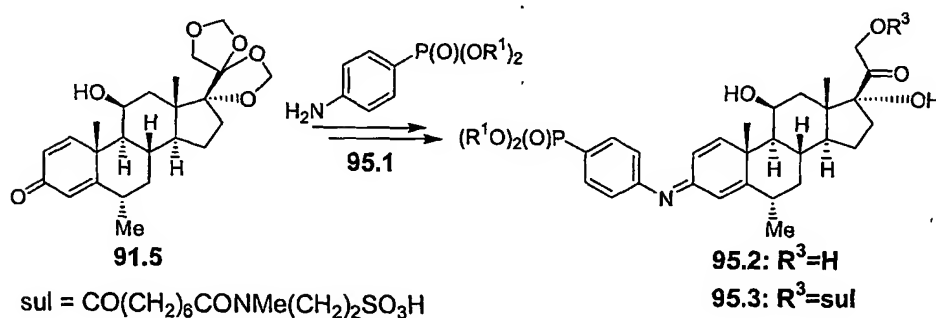
The preparation of compounds in which the phosphonate group is attached by means of a phenoxyethoxy oxime group is illustrated above. In this procedure, the dienone, 91.5, is reacted, as described above, with O-(3-bromophenoxyethyl)hydroxylamine, 94.1, prepared as described above from 3-bromophenoxyethyl bromide (FR 1,481,052) and BOC-protected hydroxylamine, to give, after deprotection of the side-chain, the oxime, 94.2. The product is then reacted, in the presence of a palladium catalyst, with a dialkyl phosphite, 94.3, to afford the phosphonate, 94.4. The preparation of arylphosphonates by means of a coupling reaction between aryl bromides and dialkyl phosphites is described in *J. Med. Chem.*, 1992, 35, 1371. The reaction is performed in an inert solvent such as toluene, in the presence of a base such as triethylamine and a catalytic amount of tetrakis(triphenylphosphine)palladium(0). The 21-hydroxy group is then converted into the 21-sulfate product, 94.5.

Alternatively, the bromo compound, 94.2, is coupled with a dialkyl propenylphosphonate, 94.6 (Aldrich), to afford the phosphonate, 94.7. The coupling of aryl halides with olefins by means of the Heck reaction is described, for example, in "Advanced Organic Chemistry," by F. A. Carey and R. J. Sundberg, Plenum, 2001, p. 503ff and in *Acc. Chem. Res.*, 1979, 12, 146. The aryl bromide and the olefin are coupled in a polar solvent such as dimethylformamide or dioxan, in the presence of a palladium(0) catalyst such as

tetrakis(triphenylphosphine)palladium(0) or palladium(II) catalyst such as palladium(II) acetate, and optionally in the presence of a base such as triethylamine or potassium carbonate. Optionally, the styrenoid double bond present in the product, **94.7**, is reduced, for example by reaction with diimide, to produce the saturated analog, **94.9**. The reduction of olefinic bonds is described in "Comprehensive Organic Transformations," by R. C. Larock, VCH, 1989, p. 6ff. The transformation is effected by means of catalytic hydrogenation, for example using a palladium on carbon catalyst and hydrogen or a hydrogen donor, or by the use of diimide or diborane. The products, **94.7** and **94.9**, are then converted into the suleptanate esters, **94.8** and **94.10**.

Using the above procedures, but employing, in place of the bromophenoxyethoxy reagent, **94.1**, different bromo-substituted aryl or heteroaryl alkoxy hydroxylamines, and/or different dialkyl alkenyl phosphonates, the products analogous to the compounds, **94.5**, **94.8** and **94.10** are obtained.

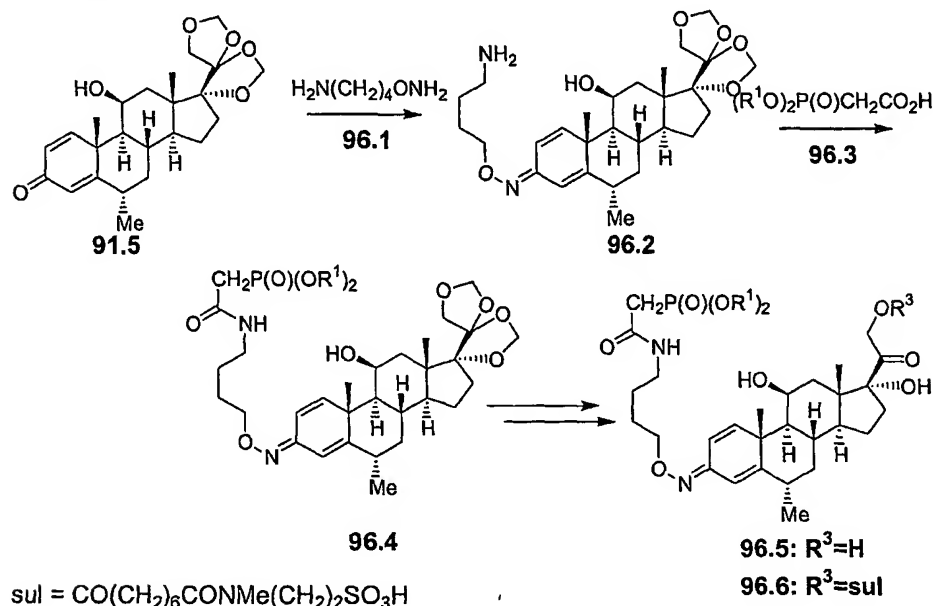
Example 95



The preparation of phosphonates in which the phosphonate is attached by means of an imino group is illustrated above. In this procedure, compound, **91.5**, is reacted with a dialkyl 4-aminophenyl phosphonate, **95.1** (Epsilon), to give, after deprotection, the imine product, **95.2**. The reaction is conducted in a hydrocarbon solvent such as toluene or xylene, at reflux temperature, in the presence of a basic catalyst such as sodium methoxide, or an acid catalyst such as p-toluenesulfonic acid, under azeotropic conditions. The product is then converted into the suleptanate ester, **95.3**.

Using the above procedures, but employing, in place of the 4-aminophenyl phosphonate, **95.1**, different amino-substituted aryl or heteroaryl phosphonates, products analogous to **95.3** are obtained.

Example 96



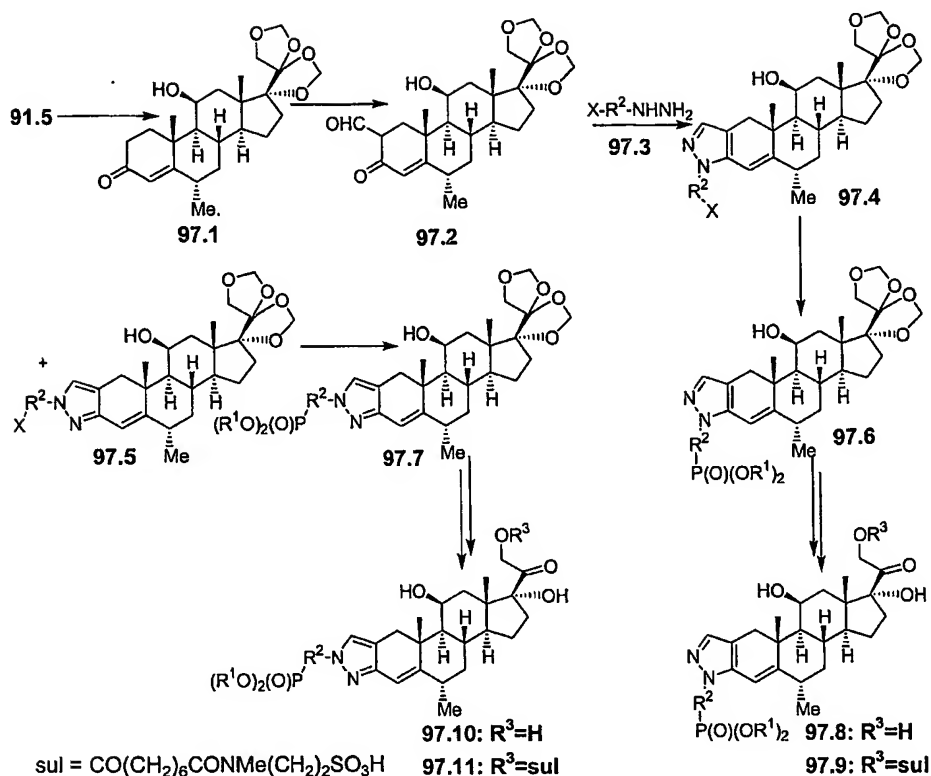
The preparation of phosphonates in which the phosphonate is attached by means of an oximino group and an amide linkage is illustrated above. In this procedure, the dienone, **91.5**, is reacted with O-(4-aminobutyl)hydroxylamine, **96.1** (*Pol. J. Chem.*, 1981, 55, 1163), to yield the oxime, **96.2**. The reaction of steroidal 1,4-dien-3-ones with substituted hydroxylamines is described in *J. Steroid Bioch.*, 1976, 7, 795. The reaction is performed between equimolar amounts of the reactants in a polar organic solvent such as pyridine or methanol, optionally in the presence of acetic acid or sodium acetate. The oxime is then coupled with a dialkyl phosphonoacetic acid, **96.3** (Aldrich), to yield the amide oxime, **96.4**. The preparation of amides from carboxylic acids and derivatives is described, for example, in "Organic Functional Group Preparations," by S.R.Sandler and W. Karo, Academic Press, 1968, p. 274, and "Comprehensive Organic Transformations," by R. C. Larock, VCH, 1989, p. 972ff. The carboxylic acid is reacted with the amine in the presence of an activating agent, such as, for example, dicyclohexylcarbodiimide or diisopropylcarbodiimide, optionally in the presence of, for example, hydroxybenztriazole, N-

hydroxysuccinimide or N-hydroxypyridone, in a non-protic solvent such as, for example, pyridine, DMF or dichloromethane, to afford the amide.

Alternatively, the carboxylic acid is first converted into an activated derivative such as the acid chloride, anhydride, mixed anhydride, imidazolidine and the like, and then reacted with the amine, in the presence of an organic base such as, for example, pyridine, to afford the amide. The conversion of a carboxylic acid into the corresponding acid chloride is effected by treatment of the carboxylic acid with a reagent such as, for example, thionyl chloride or oxalyl chloride in an inert organic solvent such as dichloromethane, optionally in the presence of a catalytic amount of dimethylformamide.

The amide product, **96.4**, is then converted, as described herein, into the suleptanate, **96.6**.

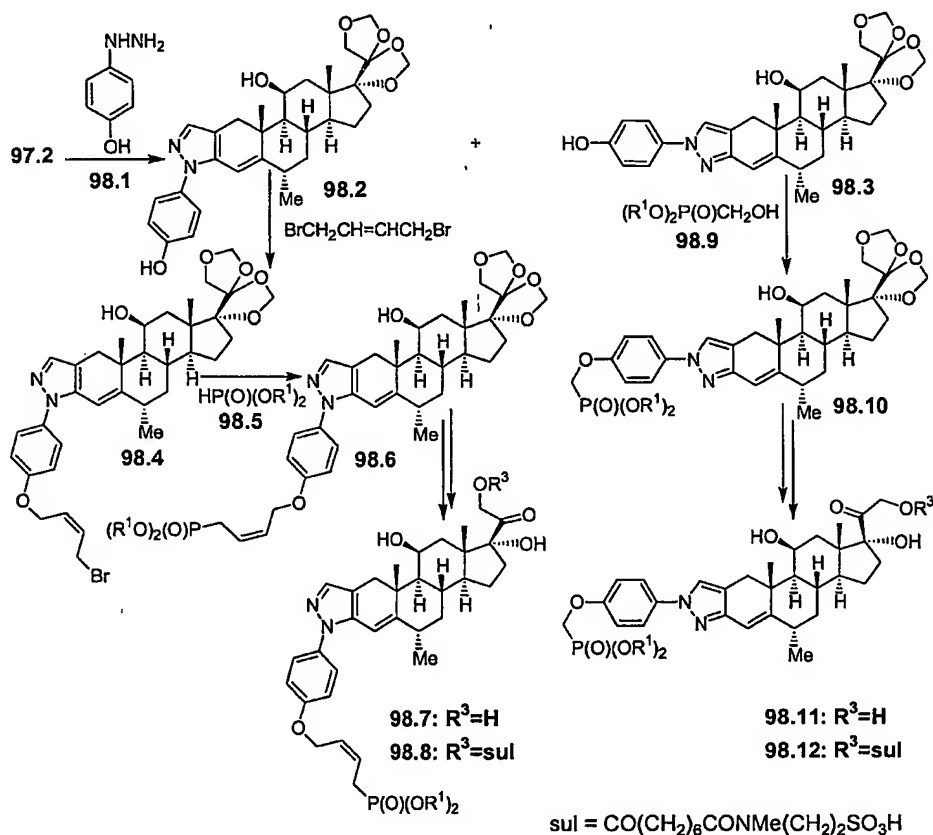
Using the above procedures, but employing, in place of the hydroxylamine, **96.1**, different amino-substituted hydroxylamines, and/or different carboxy-substituted phosphonates, the products analogous to **96.6** are obtained.

Example 97

The preparation of the phosphonate esters in which the phosphonate group is attached to the 1' or 2' position of the pyrazole ring, by means of an aromatic or heteroaromatic group, a heteroatom and a variable carbon chain is illustrated above. In this procedure, the BMD-protected dienone, **91.5**, is reduced to afford the 1,2-dihydro product, **97.1**. The catalytic hydrogenation reaction is effected by the use of tris(triphenylphosphine)rhodium (I) chloride, for example as described in *J. Med. Chem.*, **2001**, *44*, 602. The product is then reacted with ethyl formate and a base such as sodium hydride, in an inert solvent such as toluene or dimethylformamide, as described in *J. Am. Chem. Soc.*, **1964**, *86*, 1520, to afford the 2-formyl product, **97.2**. This compound is then reacted with an alkyl, aralkyl, aryl or heteroaryl hydrazine, **97.3**, wherein R^2 is alkyl, aralkyl, aryl or heteroaryl and in which the substituent X is either a phosphonate group or a group which is subsequently transformed into a phosphonate-containing substituent. For example, X is dialkylphosphono, bromo, hydroxy, amino, carboxyl and the like. The reaction yields the isomeric 2'- and 1'-aryl pyrazoles, **97.4** and **97.5**. The pyrazole-forming reaction is performed between

equimolar amounts of the reactants in an acidic solvent such as acetic acid, as described in *J. Am. Chem. Soc.*, 1964, 86, 1520. The pyrazoles, 97.4 and 97.5, are then transformed, for example by the procedures described in Examples 98 and 99, via the BMD-protected intermediates, 97.6 and 97.7, into the phosphonate suleptanates, 97.9 and 97.11.

Example 98

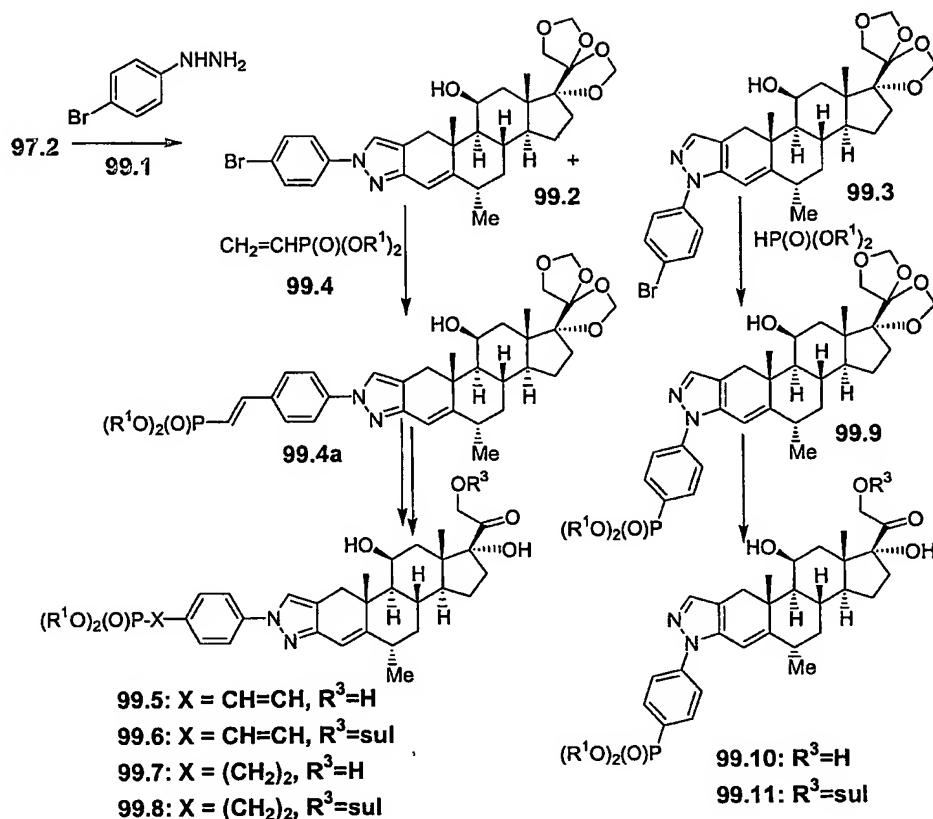


The preparation of phosphonates in which the phosphonate is attached by means of a phenyl ring and an alkoxy or an alkenyl linkage is illustrated above. In this procedure, the ketoaldehyde, 97.2, is reacted, as described above, with 4-hydroxyphenylhydrazine, 98.1 (Epsilon), to give the pyrazoles, 98.2 and 98.3. The 2'-substituted isomer, 98.2, is then reacted in dimethylformamide solution at ambient temperature with one molar equivalent of 1,4-dibromobut-2-ene and dimethylaminopyridine, to yield the bromoether, 98.4. The product is then reacted at 120° in an Arbuzov reaction with a trialkyl phosphite, 98.5, to give the phosphonate product, 98.6. The Arbuzov reaction, in which an alkyl bromide is

transformed into the corresponding phosphonate, by heating at from 60° to about 150° with a trialkyl phosphite, is described in *Handb. Organophosphorus Chem.*, 1992, 115-72. The BMD protecting group is then removed and the product is acylated to yield the suleptanate ester triol, 98.8.

Alternatively, the 1'-substituted pyrazole, 98.3, is reacted, in a Mitsunobu reaction, with a dialkyl 2-hydroxymethyl phosphonate, 98.9 (Aldrich), to afford the ether, 98.10. The preparation of aromatic ethers by means of the Mitsunobu reaction is described, for example, in "Comprehensive Organic Transformations," by R. C. Larock, VCH, 1989, p. 448, and in "Advanced Organic Chemistry," Part B, by F.A. Carey and R. J. Sundberg, Plenum, 2001, p. 153-4 and in *Org. React.*, 1992, 42, 335. The phenol and the alcohol or thiol component are reacted together in an aprotic solvent such as, for example, tetrahydrofuran, in the presence of a dialkyl azodicarboxylate and a triarylphosphine, to afford the ether or thioether products. The procedure is also described in *Org. React.*, 1992, 42, 335. The product, 98.10, is then deprotected to give the triol, 98.11, and the latter compound is acylated to afford the suleptanate, 98.12.

Using the above procedures, but employing different dibromides or hydroxyl-substituted phosphonates, the products analogous to 98.8 and 98.12 are obtained. The functionalization procedures are interchangeable between the pyrazole substrates, 98.2 and 98.3.

Example 99

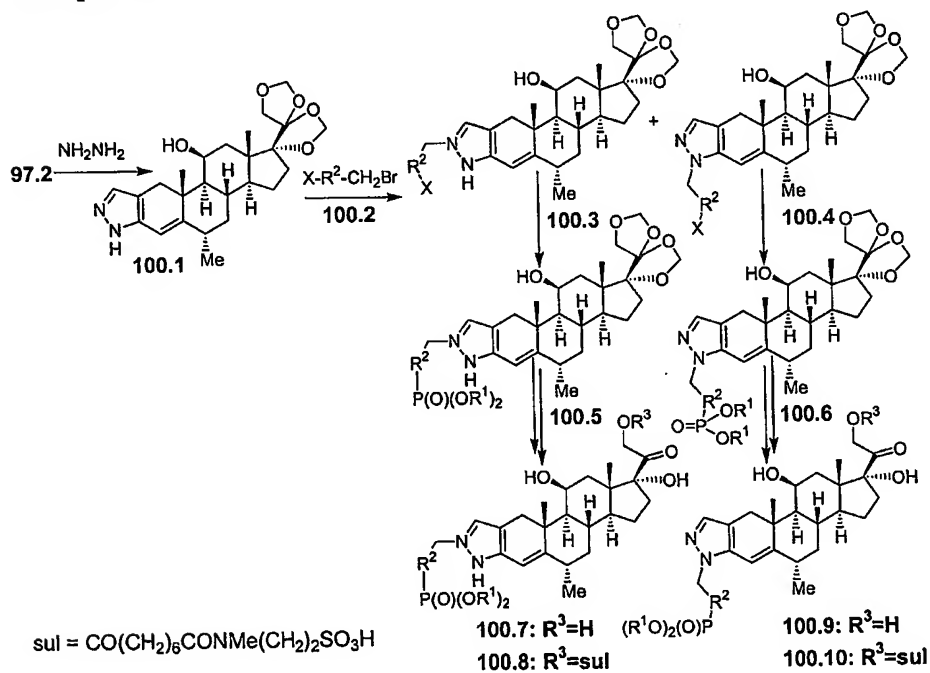
The preparation of the phosphonates in which the phosphonate group is attached by means of a phenyl ring or a phenyl ring and a saturated or unsaturated carbon chain is illustrated above. In this procedure, the keto-aldehyde, 97.2, is reacted, as described above, with 4-bromophenyl hydrazine, 99.1 (*J. Organomet. Chem.*, 1999, 62, 581), to produce the pyrazoles, 99.2 and 99.3. The 1'-substituted isomer, 99.2, is coupled, in the presence of a palladium catalyst, with a dialkyl vinylphosphonate, 99.4 (Aldrich), to give the phosphonate, 99.4a. The product is then deprotected to afford the triol, 99.5, which is converted into the sulfate, 99.6. Optionally, the styrenoid double bond present in the product, 99.6, is reduced, as described above, to produce the saturated analog, 99.8.

Alternatively, the 2'-substituted pyrazole, 99.3, is coupled, in the presence of a palladium catalyst, with a dialkyl phosphite to prepare the phosphonate, 99.9, which is deprotected, and the product is acylated to give the

suleptanate ester, **99.11**. The preparation of arylphosphonates by means of a coupling reaction between aryl bromides and dialkyl phosphites is described in *J. Med. Chem.*, **1992**, *35*, 1371. This reaction is performed in an inert solvent such as toluene, in the presence of a base such as triethylamine and tetrakis-(triphenylphosphine)palladium(0).

Using the above procedures, but employing, in place of the bromophenyl hydrazine, **99.1**, different bromo-substituted aralkyl, aryl or heteroaryl alkoxy hydrazines, and/or different dialkyl alkenyl phosphonates, the products analogous to the compounds, **99.6**, **99.8** and **99.11** are obtained.

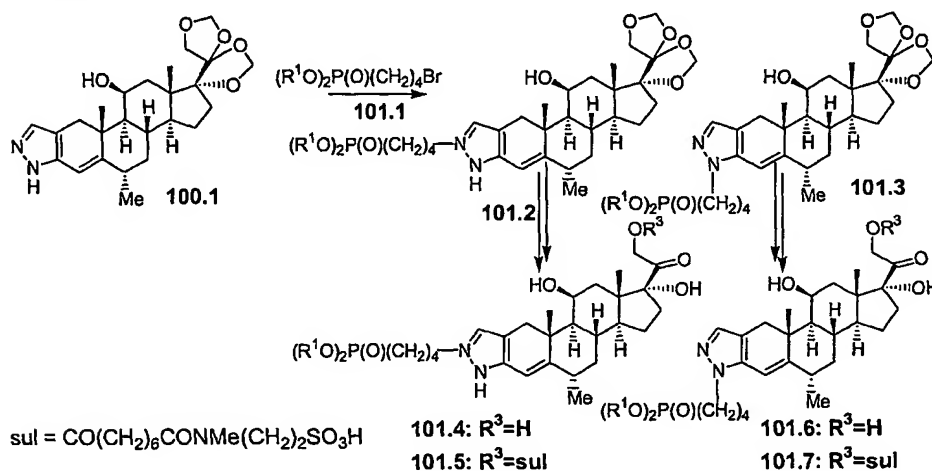
Example 100



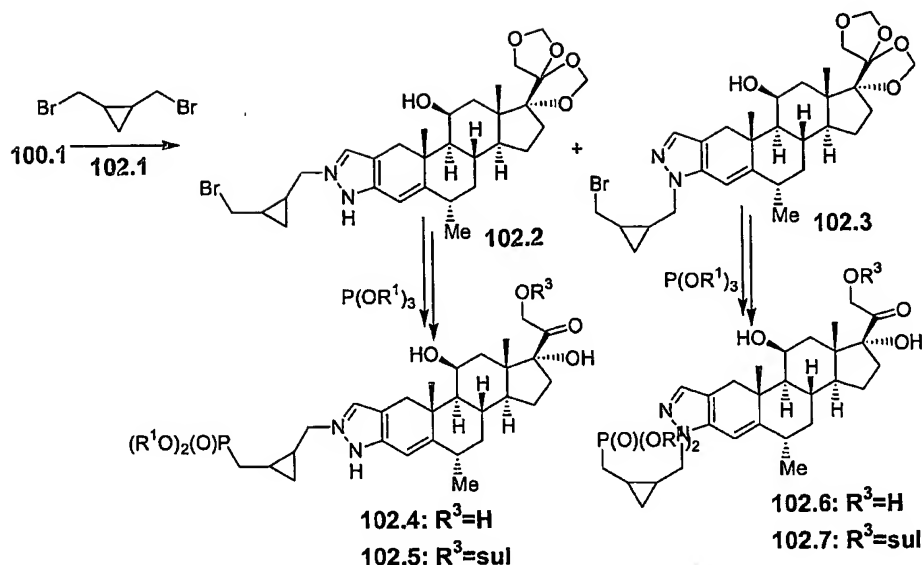
The preparation of the phosphonate esters in which the phosphonate group is attached by means of a variable carbon linkage is illustrated above. In this procedure, the ketoaldehyde, **97.2**, is reacted with hydrazine, to afford the pyrazole derivative, **100.1**. The reaction of steroidal 2-formyl-3-ketones with hydrazine is described in *J. Am. Chem. Soc.*, **1964**, *86*, 1520. The reaction is performed in acetic acid at ambient temperature. The pyrazole product is then reacted with a bromomethyl compound, **100.2**, in which R^2 and X are as defined above, to yield the alkylation products, **100.3** and **100.4**. The alkylation of

substituted pyrazoles is described, for example, in "Heterocyclic Chemistry," by T. L. Gilchrist, Longman, 1992, p. 309. The reaction is performed between equimolar amounts of the substrates in a polar solvent such as dimethylformamide or tetrahydrofuran, in the presence of a base such as dimethylaminopyridine, lithium hexamethyldisilazide and the like. The products, 100.3 and 100.4 are, except in cases where X is dialkylphosphono, converted into the phosphonates, 100.5 and 100.6, using the procedures described herein, and deprotection/acylation then affords the suleptanate esters, 100.8 and 100.10.

Example 101



The pyrazole, 100.1, is reacted in tetrahydrofuran solution, as described above, with one molar equivalent of a dialkyl bromobutyl phosphonate, 101.1, (*Synthesis*, 1994, 9, 909) and lithium hexamethyldisilazide to give the alkylated pyrazoles, 101.2 and 101.3. Deprotection/acylation then yields the suleptanates, 101.5 and 101.7.

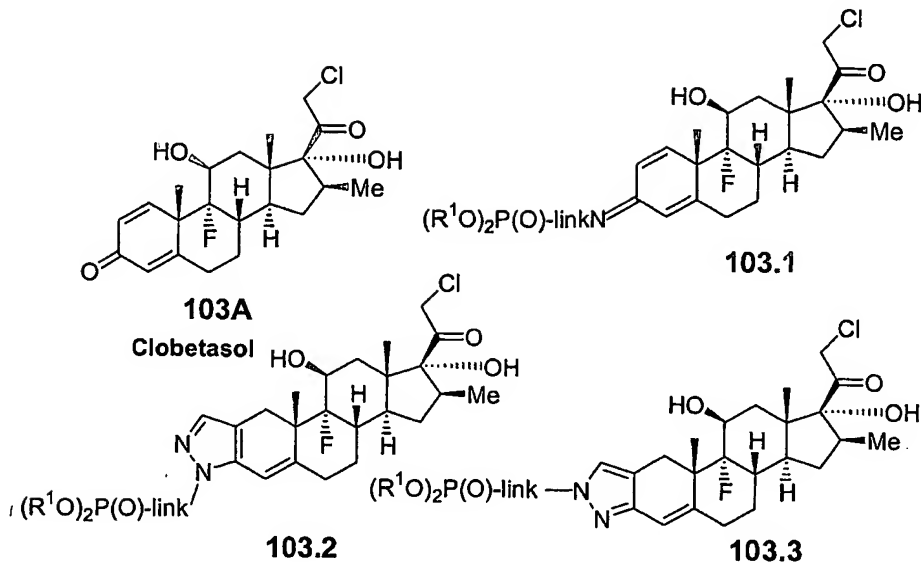
Example 102

sul = CO(CH₂)₆CONMe(CH₂)₂SO₃H

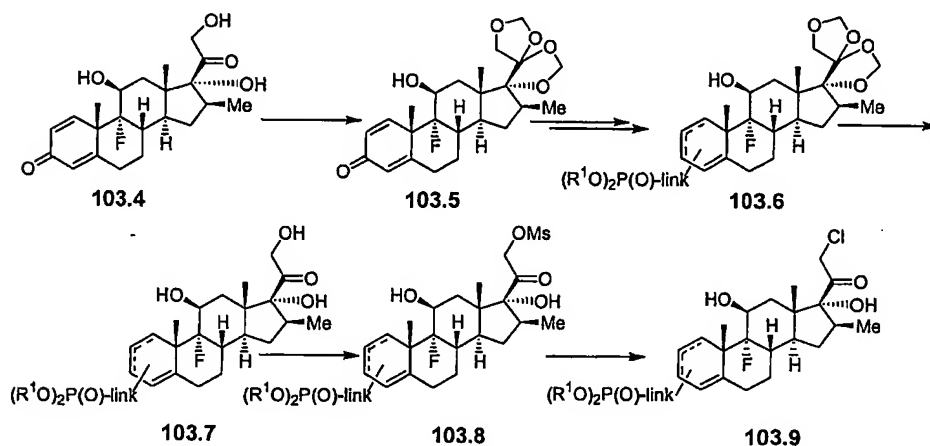
The pyrazole, **100.1**, is reacted in tetrahydrofuran solution, as described above, with 1,2-bis(bromomethyl)cyclopropane, **102.1** (*Tet.*, **1997**, 53, 10459), to give the pyrazoles, **102.2** and **102.3**. The products are subjected to an Arbuzov reaction, in which the bromomethyl substituent is converted into the dialkyl phosphonomethyl substituent, by reaction with a trialkyl phosphite at 120°, to prepare, after deprotection of the side chain and acylation, the suleptanate phosphonates, **102.5** and **102.7**. The Arbuzov reaction is described in *Handb. Organophosphorus Chem.*, **1992**, 115-72. In the procedure, the substrate is heated at from 60° to about 160° with a five to fifty-fold molar excess of the trialkyl phosphite.

Using the above procedures, but employing, in place of the dibromide, **102.1**, different dibromides, the products analogous to **102.5** and **102.7** are obtained.

Example 103



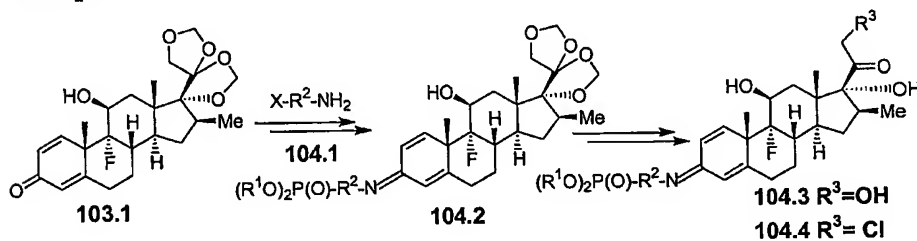
The structures of Clobetasol, 103A (U.S. Patent No. 3,721,687), and the phosphonate esters, 103.1 – 103.3, are shown above. The compounds, 103.1 – 103.3, incorporate a phosphonate moiety $(R^1O)_2P(O)$ connected to the nucleus by means of a variable linking group, designated as "link" in the attached structures.



A protection-deprotection sequence in which the steroid side-chain is protected as a bis-methylenedioxy (BMD) moiety is illustrated above. In this sequence, 9α-fluoro-16β-methyl-11β,17α,21-trihydroxypregn-1,4-dien-3,21-dione, 103.4 (U.S. Patent No. 3,721,687), is reacted with paraformaldehyde and an acid catalyst such as hydrochloric acid, as described in "Protective Groups in

Organic Synthesis," by T.W. Greene and P.G.M. Wuts, Wiley, Second Edition 1990, p. 223, to yield the BMD derivative, 103.5. The phosphonate moiety is then introduced, using the procedures described below, to produce the phosphonate ester, 103.6. The BMD moiety is then hydrolyzed, for example by treatment with 50% aqueous acetic acid, as described in "Protective Groups in Organic Synthesis," by T.W. Greene and P.G.M. Wuts, Wiley, Second Edition 1990, p. 223, to afford the triol, 103.7. The 21-hydroxy group is then converted into the 21-chloro group as described in U.S. Patent No. 3,721,687, *Chimia*, 1992, 46, 338, or *J. Med. Chem.*, 1987, 30, 1581. In this procedure, the 21-hydroxy substrate is reacted at about 0° with one molar equivalent of methanesulfonyl chloride in a basic solvent such as pyridine, to afford the 21-mesylate, 103.8. The product is then reacted, in dimethylformamide solution at about 70°, with ca. five molar equivalents of lithium chloride, to yield the 21-chloro product, 103.9.

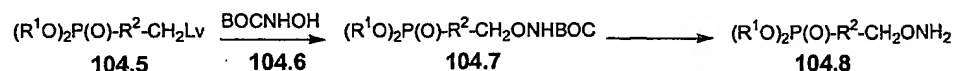
Example 104



The the preparation of phosphonates in which the phosphonate is attached by means of an imino or iminoxy group and a variable carbon chain is described herein. In this procedure, the BMD-protected derivative, 103.1, is reacted with an amine or hydroxylamine, 104.1, in which R² is an alkyl, alkenyl, cycloalkyl or cycloalkenyl group, optionally incorporating a heteroatom O, S or N, or a functional group such as an amide, ester, oxime, sulfoxide or sulfone etc, or an optionally substituted aryl, heteroaryl or aralkyl group, optionally incorporating a heteroatom O, S or N, and X is either a phosphonate group or a group which is subsequently converted into a phosphonate-containing substituent. For example, X is dialkylphosphono, bromo, hydroxy, amino, carboxy and the like. The reaction is conducted between equimolar amounts of the reactants in an aprotic solvent such as pyridine or xylene, or in an alcoholic

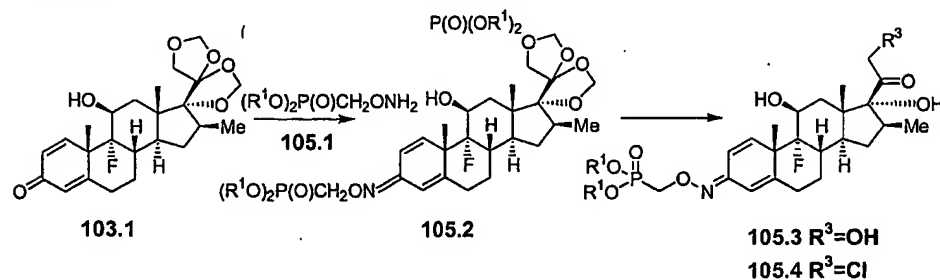
solvent such as ethanol, optionally in the presence of an acid catalyst, to give the imine or oxime. The preparation of oximes of steroidal 3-ketones is described in *Anal. Bioch.*, 1978, 86, 133 and in *J. Mass. Spectrom.*, 1995, 30, 497. The BMD-protected side-chain compound, 104.2, is then converted into the triol, 104.3, and then to the 21-chloro product, 104.4, as described herein.

Example 104A



The preparation of hydroxylamine ethers incorporating a phosphonate group is illustrated above. In this procedure, a phosphonate, 104.5, in which Lv is a leaving group such as bromo or trifluoromethylsulfonyloxy, is reacted with BOC-hydroxylamine, 104.6 (Aldrich), to produce the ether, 104.7. The reaction is conducted between equimolar amounts of the reactants in a polar solvent such as dimethylformamide or tetrahydrofuran, in the presence of a base such as potassium hydroxide or dimethylaminopyridine. Deprotection, for example by treatment with trifluoroacetic acid, then gives the hydroxylamine ether, 104.8.

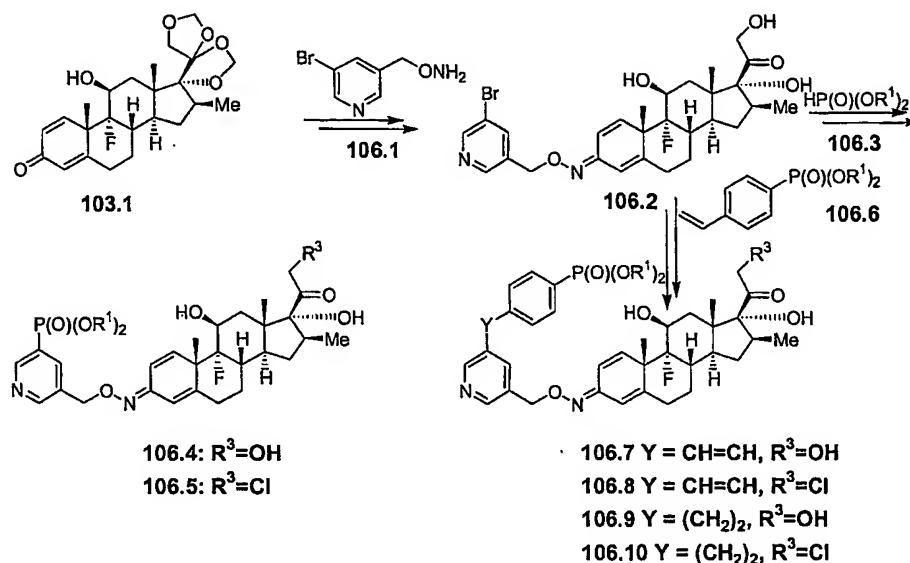
Example 105



The preparation of phosphonates in which the phosphonate is attached by means of an iminoxy group. In this procedure, the substrate, 103.1, is reacted with a dialkyl phosphonomethyl hydroxylamine, 105.1, prepared as described above from a dialkyl trifluoromethylsulfonyloxymethyl phosphonate (*Tetrahedron Lett.*, 1986, 27, 1477) and BOC-hydroxylamine, to afford the oxime, 105.2. Deprotection then affords the triol, 105.3, from which the 21-chloro compound, 105.4, is prepared. The oxime forming reaction is performed at ambient temperature in ethanol-acetic acid solution between equimolar amounts of the reactants. Using the above procedures, but employing, in place

of the hydroxylamine ether, **105.1**, different oxime ethers, **104.1**, the corresponding products, **105.4** are obtained.

Example 106

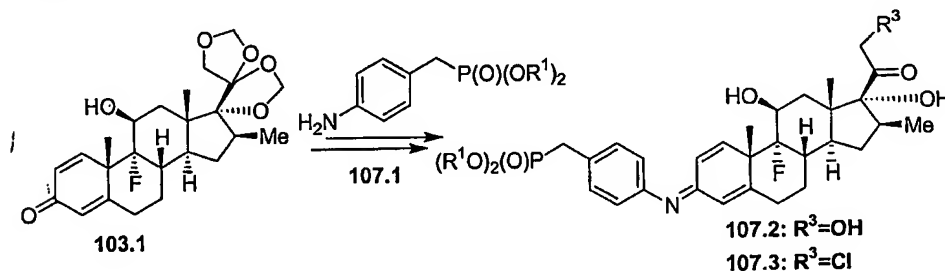


The preparation of compounds in which the phosphonate group is attached by means of a 3-pyridylmethoxy oxime group described herein. In this procedure, the dienone, **103.1**, is reacted, as described above, with O-(5-bromo-3-pyridylmethoxy)hydroxylamine, **106.1**, prepared as described above from 5-bromo-3-bromomethylpyridine (WO 95/28400) and BOC-protected hydroxylamine, to give, after deprotection of the side-chain, the oxime, **106.2**. The product is then reacted, in the presence of a palladium catalyst, with a dialkyl phosphite, **106.3**, to afford the phosphonate, **106.4**. The preparation of arylphosphonates by means of a coupling reaction between aryl bromides and dialkyl phosphites is described in *J. Med. Chem.*, **1992**, *35*, 1371. The reaction is performed in an inert solvent such as toluene, in the presence of a base such as triethylamine and a catalytic amount of tetrakis(triphenylphosphine)-palladium(0). The 21-hydroxy group is then converted into the 21-chloro derivative, **106.5**.

Alternatively, the bromo compound, **106.2**, is coupled with a dialkyl 4-vinylphenyl phosphonate, **106.6** (*Macromolecules*, **1998**, *31*, 2918), to afford the phosphonate, **106.7**. The coupling of aryl halides with olefins by means of the Heck reaction is described, for example, in "Advanced Organic Chemistry," by

F. A. Carey and R. J. Sundberg, Plenum, 2001, p. 503ff and in *Acc. Chem. Res.*, 1979, 12, 146. The aryl bromide and the olefin are coupled in a polar solvent such as dimethylformamide or dioxan, in the presence of a palladium(0) catalyst such as tetrakis(triphenylphosphine)palladium(0) or palladium(II) catalyst such as palladium(II) acetate, and optionally in the presence of a base such as triethylamine or potassium carbonate. Optionally, the styrenoid double bond present in the product, 106.7, is reduced, for example by reaction with diimide, to produce the saturated analog, 106.9. The reduction of olefinic bonds is described in "Comprehensive Organic Transformations," by R. C. Larock, VCH, 1989, p. 6ff. The transformation is effected by means of catalytic hydrogenation, for example using a palladium on carbon catalyst and hydrogen or a hydrogen donor, or by the use of diimide or diborane. The products, 106.7 and 106.9, are then converted into the 21-chloro analogs, 106.8 and 106.10. Using the above procedures, but employing, in place of the bromopyridyl-methoxy reagent, 106.1, different bromo-substituted aryl or heteroaryl alkoxy hydroxylamines, and/or different dialkyl alkenyl phosphonates, the products analogous to the compounds, 106.5, 106.8 and 106.10 are obtained.

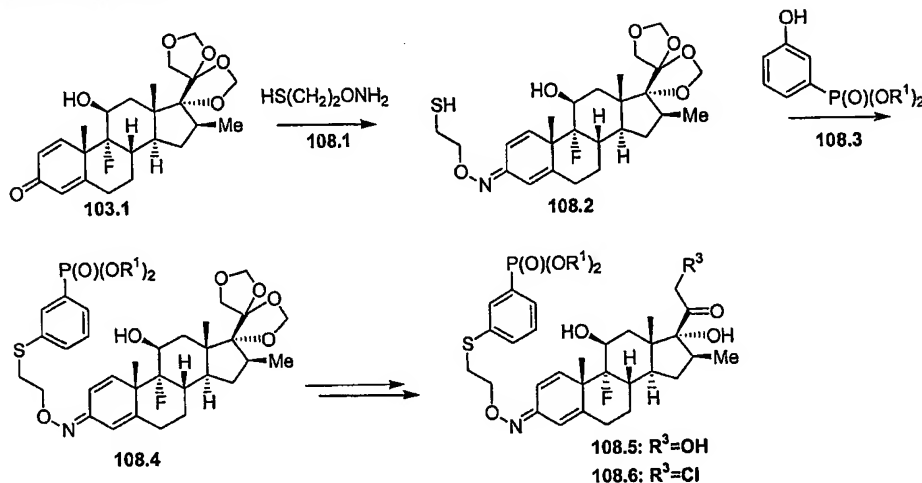
Example 107



The preparation of phosphonates in which the phosphonate is attached by means of an imino group is illustrated above. In this procedure, the substrate, 103.1, is reacted with a dialkyl 4-aminobenzyl phosphonate, 107.1, (Fluka) to give, after deprotection, the imine product, 107.2. The reaction is conducted in a hydrocarbon solvent such as toluene or xylene, at reflux temperature, in the presence of a basic catalyst such as sodium methoxide, or an acid catalyst such as p-toluenesulfonic acid, under azeotropic conditions. The product is then converted into the 21-chloro compound, 107.3.

Using the above procedures, but employing, in place of the 4-aminobenzyl phosphonate, 107.1, different amino-substituted aryl or heteroaryl phosphonates, products analogous to 107.3 are obtained.

Example 103

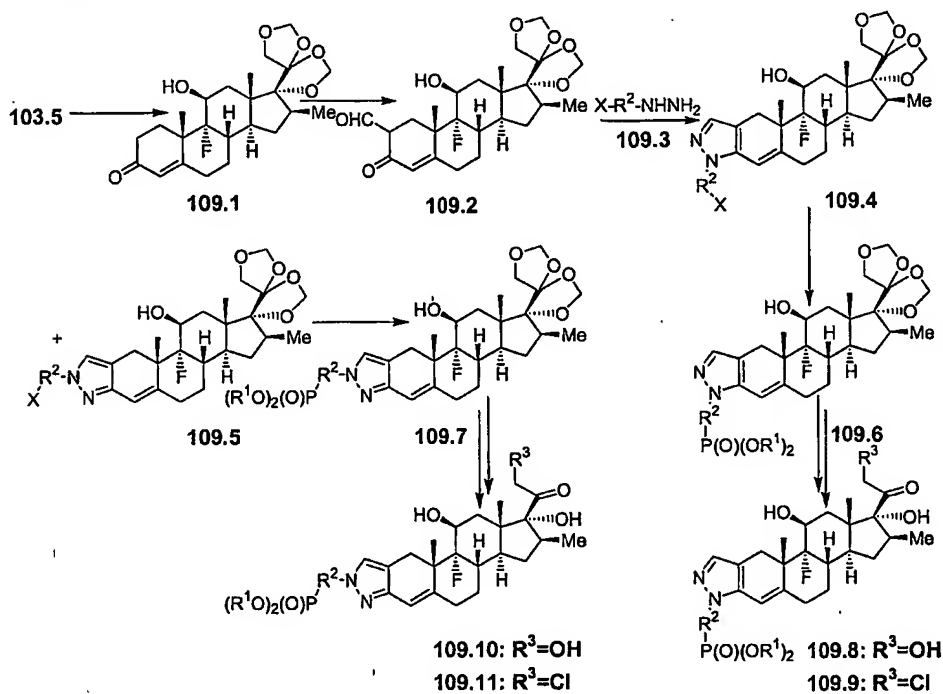


The preparation of phosphonates in which the phosphonate is attached by means of an oximino group and a thioether linkage is illustrated above. In this procedure, the dienone, 103.1, is reacted with O-(2-mercaptoethyl)hydroxylamine, 108.1 (*Bioorganicheskaya Khim.*, 1986, 12, 1662), to yield the oxime, 108.2. The reaction of steroidal 1,4-dien-3-ones with substituted hydroxylamines is described in *J. Steroid Bioch.*, 1976, 7, 795; the reaction is performed between equimolar amounts of the reactants in a polar organic solvent such as pyridine or methanol, optionally in the presence of acetic acid or sodium acetate. The product is then coupled, in a Mitsunobu reaction, with a dialkyl 3-hydroxyphenyl phosphonate, 108.3 (Aurora), to yield the thioether oxime, 108.4. The preparation of aromatic ethers by means of the Mitsunobu reaction is described, for example, in "Comprehensive Organic Transformations", by R. C. Larock, VCH, 1989, p. 448, and in "Advanced Organic Chemistry," Part B, by F.A. Carey and R. J. Sundberg, Plenum, 2001, p. 153-4 and in *Org. React.*, 1992, 42, 335. The phenol and the alcohol or thiol component are reacted together in an aprotic solvent such as, for example, tetrahydrofuran, in the presence of a dialkyl azodicarboxylate and a triarylphosphine, to afford the ether or thioether products. The procedure is also described in *Org. React.*, 1992, 42, 335-656.

The thioether product, **108.4**, is then hydrolyzed and converted into the 21-chloro product, **108.6**, as described in Example 103.

Using the above procedures, but employing, in place of the hydroxylamine, **108.3**, different hydroxy or mercapto-substituted hydroxylamines, and/or different hydroxyaryl-substituted phosphonates, the products analogous to **108.6** are obtained.

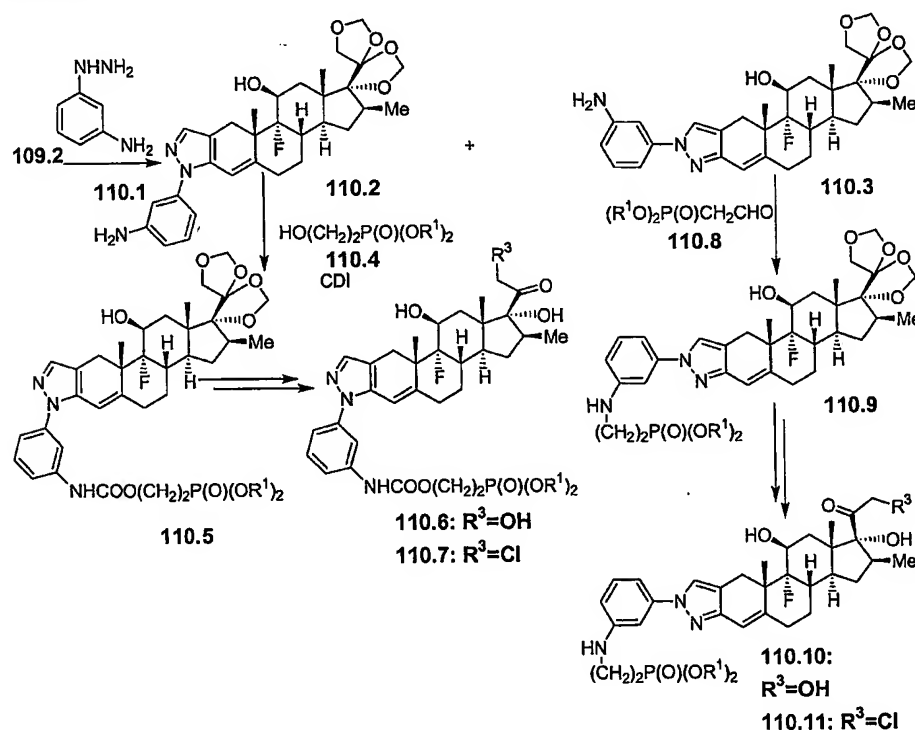
Example 109



The preparation of the phosphonate esters in which the phosphonate group is attached to the 1' or 2' position of the pyrazole ring, by means of an aromatic or heteroaromatic group, a heteroatom and a variable carbon chain is illustrated above. In this procedure, the BMD-protected dienone, **103.5**, is reduced to afford the 1,2-dihydro product, **109.1**. The catalytic hydrogenation reaction is effected by the use of tris(triphenylphosphine)rhodium (I) chloride, for example as described in *J. Med. Chem.*, 2001, 44, 602. The product is then reacted with ethyl formate and a base such as sodium hydride, in an inert solvent such as toluene or dimethylformamide, as described in *J. Am. Chem. Soc.*, 1964, 86, 1520, to afford the 2-formyl product, **109.2**. This compound is then reacted with an alkyl, aralkyl, aryl or heteroaryl hydrazine, **109.3**, wherein R^2 is alkyl,

aralkyl, aryl or heteroaryl and in which the substituent X is either a phosphonate group or a group which is subsequently transformed into a phosphonate-containing substituent. For example, X is dialkylphosphono, bromo, hydroxy, amino, carboxyl and the like. The reaction yields the isomeric 2'- and 1'-aryl pyrazoles, 109.4 and 109.5. The pyrazole-forming reaction is performed between equimolar amounts of the reactants in an acidic solvent such as acetic acid, as described in *J. Am. Chem. Soc.*, 1964, 86, 1520. The pyrazoles, 109.4 and 109.5, are then transformed, for example by the procedures described in Examples 110 and 111, via the BMD-protected intermediates, 109.6 and 109.7, into the 21-chloro phosphonates, 109.9 and 109.11.

Example 110

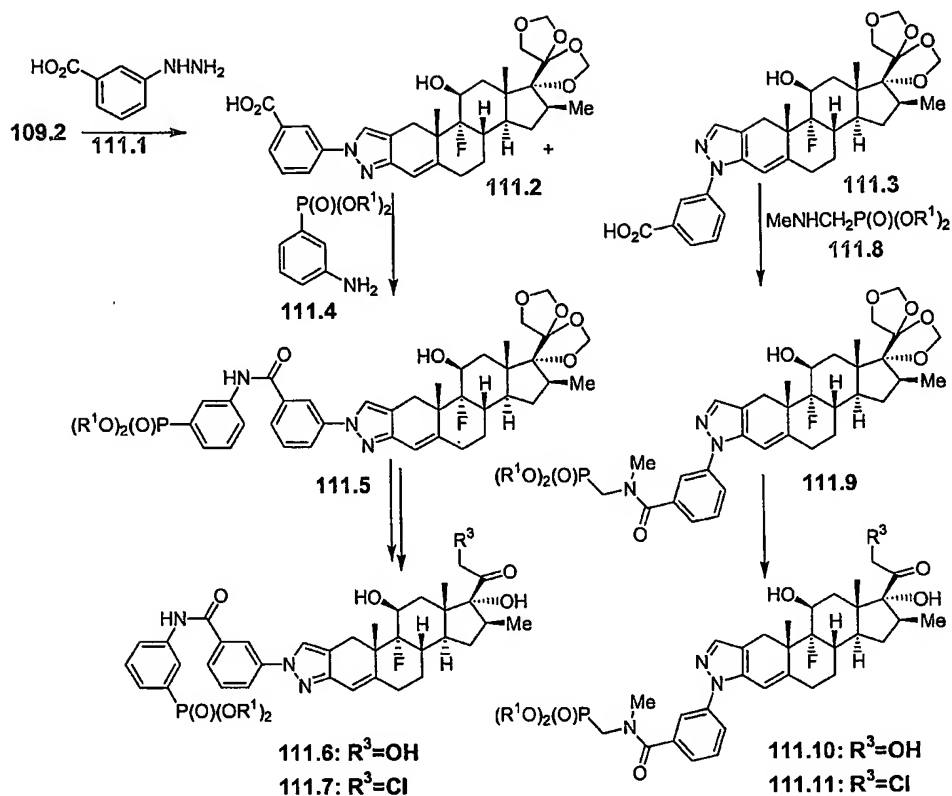


The preparation of the phosphonate esters in which the phosphonate group is attached to the 1' or 2' position of the pyrazole ring, by means of a carbamate or an amino linkage is illustrated above. In this procedure, the ketoaldehyde, 109.2, is reacted, as described above, with 3-aminophenylhydrazine 110.1 (EP 437105) to give the pyrazoles 110.2 and 110.3. The 2'-substituted isomer 110.2 is then reacted in dimethylformamide solution at

ambient temperature with one molar equivalent of a dialkyl 2-hydroxyethyl phosphonate, 110.4 (Epsilon), and carbonyl diimidazole, to yield the carbamate, 110.5. The preparation of carbamates is described in "Comprehensive Organic Functional Group Transformations," A. R. Katritzky, ed., Pergamon, 1995, Vol. 6, p 416ff, and in "Organic Functional Group Preparations," by S. R. Sandler and W. Karo, Academic Press, 1986, p. 260ff. In the procedure, the amine is reacted in an inert aprotic solvent such as dichloromethane or tetrahydrofuran, with phosgene or a functional equivalent thereof, such as carbonyl diimidazole, triphosgene, pentafluorophenyl carbonate and the like, to afford the corresponding activated acylamine. The latter compound is then reacted with an alcohol to yield the carbamate. The BMD protecting group is then removed and the product is converted into the 21-chloro product, 110.7.

Alternatively, the 1'-substituted pyrazole, 110.3, is reacted, in a reductive amination reaction, with a dialkyl formylmethyl phosphonate, 110.8 (*Zh. Obschei. Khim.*, 1987, 57, 2793), and sodium triacetoxymethylborohydride, to afford the amine, 110.9. The preparation of amines by means of reductive amination procedures is described, for example, in "Comprehensive Organic Transformations," by R. C. Larock, VCH, p 421, and in "Advanced Organic Chemistry," Part B, by F.A. Carey and R. J. Sundberg, Plenum, 2001, p 269. In this procedure, the amine component and the aldehyde or ketone component are reacted together in the presence of a reducing agent such as, for example, borane, sodium cyanoborohydride, sodium triacetoxymethylborohydride or diisobutylaluminum hydride, optionally in the presence of a Lewis acid, such as titanium tetrakisopropoxide, as described in *J. Org. Chem.*, 1990, 55, 2552. The product, 110.9, is then deprotected to give the triol, 110.10, and the latter compound is transformed into the 21-chloro analog, 110.11.

Using the above procedures, but employing different formyl or hydroxyl-substituted phosphonates, and/or different amino-substituted hydrazines, the products analogous to 110.7 and 110.11 are obtained. The functionalization procedures are interchangeable between the pyrazole substrates, 110.2 and 110.3.

Example 111

The preparation of the phosphonates in which the phosphonate group is attached by means of a phenyl ring and an amide linkage is illustrated above. In this procedure, the ketoaldehyde, **109.2**, is reacted, as described above, with 3-carboxyphenyl hydrazine, **111.1** (Apin), to produce the pyrazoles, **111.2** and **111.3**. The 1'-substituted isomer, **111.1**, is coupled, in the presence of dicyclohexylcarbodiimide, with a dialkyl 3-aminophenyl phosphonate, **111.4** (Aurora), to give the amide, **111.5**. The preparation of amides from carboxylic acids and derivatives is described, for example, in "Organic Functional Group Preparations," by S.R.Sandler and W. Karo, Academic Press, 1968, p. 274, and "Comprehensive Organic Transformations," by R. C. Larock, VCH, 1989, p. 972ff. The carboxylic acid is reacted with the amine in the presence of an activating agent, such as, for example, dicyclohexylcarbodiimide or diisopropylcarbodiimide, optionally in the presence of, for example, hydroxybenztriazole, N-hydroxysuccinimide or N-hydroxypyridone, in a non-

protic solvent such as, for example, pyridine, DMF or dichloromethane, to afford the amide.

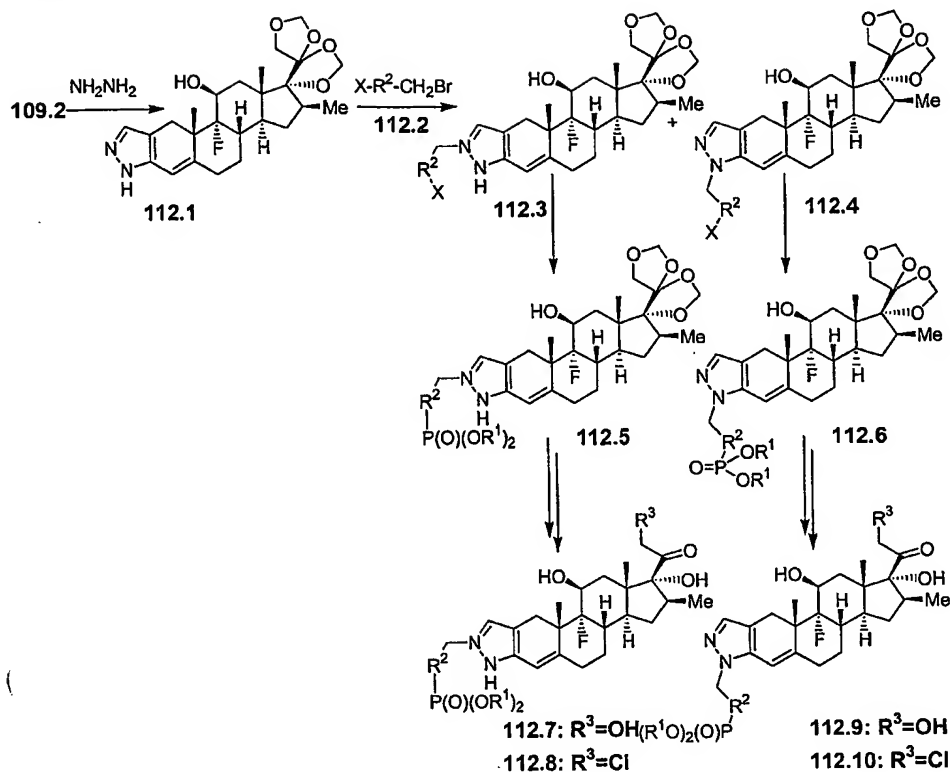
Alternatively, the carboxylic acid may first be converted into an activated derivative such as the acid chloride, anhydride, mixed anhydride, imidazolid and the like, and then reacted with the amine, in the presence of an organic base such as, for example, pyridine, to afford the amide.

The conversion of a carboxylic acid into the corresponding acid chloride can be effected by treatment of the carboxylic acid with a reagent such as, for example, thionyl chloride or oxalyl chloride in an inert organic solvent such as dichloromethane, optionally in the presence of a catalytic amount of dimethylformamide.

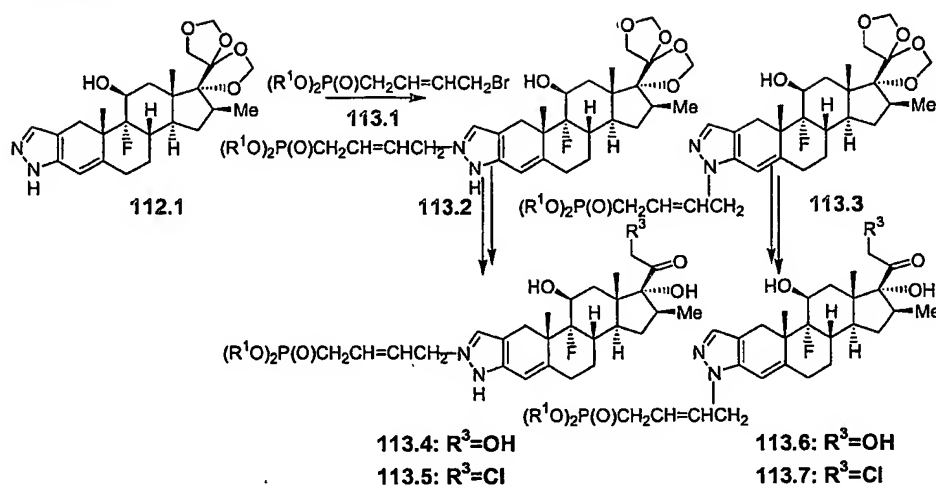
The product is then deprotected to afford the triol, 111.6, which is converted into the 21-chloro compound, 111.7.

Alternatively, the 2'-substituted pyrazole, 111.3, is coupled, as described above, with a dialkyl methylaminomethyl phosphonate, 111.8, to prepare the amide phosphonate 111.9 which is deprotected, and the product is converted into the 21-chloro analog, 111.11.

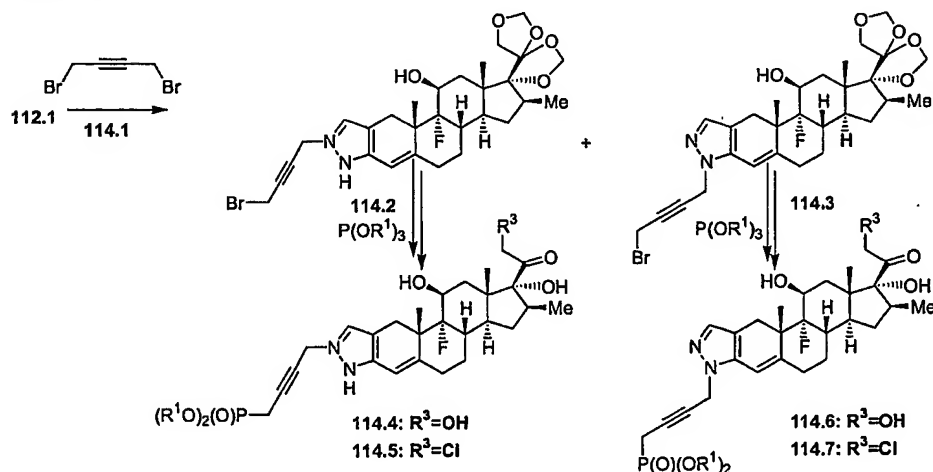
Using the above procedures, but employing, in place of the carboxyphenyl hydrazine, 111.1, different carboxy-substituted aralkyl, aryl or heteroaryl alkoxy hydrazines, and/or different dialkyl amino-substituted phosphonates, the products analogous to the compounds, 111.7 and 111.11 are obtained.

Example 112

The preparation of the phosphonate esters in which the phosphonate group is attached by means of a variable carbon linkage is illustrated above. In this procedure, the ketoaldehyde, **109.2**, is reacted with hydrazine, to afford the pyrazole derivative, **112.1**. The reaction of steroidal 2-formyl-3-ketones with hydrazine is described in *J. Am. Chem. Soc.*, **1964**, 86, 1520. The reaction is performed in acetic acid at ambient temperature. The pyrazole product is then reacted with a bromomethyl compound, **112.2**, in which R^2 and X are as defined above, to yield the alkylation products, **112.3** and **112.4**. The alkylation of substituted pyrazoles is described, for example, in "Heterocyclic Chemistry," by T. L. Gilchrist, Longman, **1992**, p. 309. The reaction is performed between equimolar amounts of the substrates in a polar solvent such as dimethylformamide or tetrahydrofuran, in the presence of a base such as dimethylaminopyridine, lithium hexamethyldisilazide and the like. The products, **112.3** and **112.4**, are, except in cases where X is dialkylphosphono, converted into the phosphonates, **112.5** and **112.6**, using the procedures described herein, and deprotection/acylation then affords the 21-chloro compounds, **112.8** and **112.10**.

Example 113

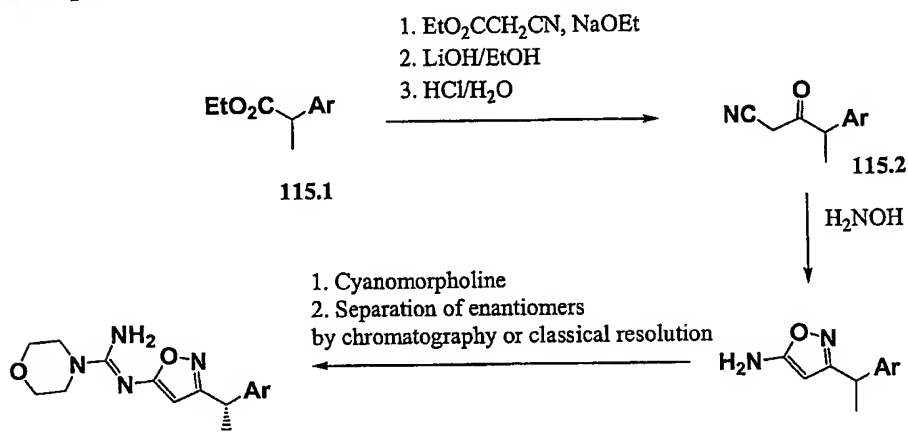
The pyrazole, **112.1**, is reacted in tetrahydrofuran solution, as described above, with one molar equivalent of a dialkyl bromobutenyl phosphonate, **113.1** (*J. Med. Chem.*, **1992**, 35, 1371), and lithium hexamethyldisilazide to give the alkylated pyrazoles, **113.2** and **113.3**. Deprotection followed by chlorination yields the 21-chloro products, **113.5** and **113.7**.

Example 114

The pyrazole, **112.1**, is reacted in tetrahydrofuran solution, as described above, with 1,4-dibromobut-2-yne, **114.1** (Aldrich), to give the pyrazoles, **114.2** and **114.3**. The products are subjected to an Arbuzov reaction, in which the bromomethyl substituent is converted into the dialkyl phosphonomethyl substituent, by reaction with a trialkyl phosphite at 120°, to prepare, after

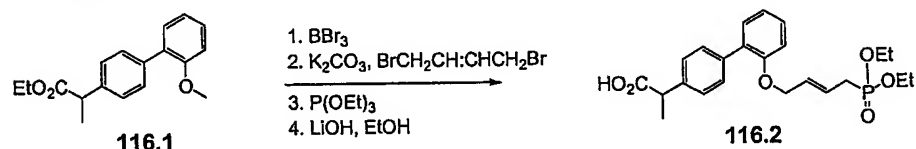
deprotection of the side chain and chlorination, the 21-chloro phosphonates, **114.5** and **114.7**. The Arbuzov reaction is described in *Handb. Organophosphorus Chem.*, 1992, 115-72. In the procedure, the substrate is heated at from 60° to about 160° with a five to fifty-fold molar excess of the trialkyl phosphite. Using the above procedures, but employing, in place of the dibromide, **114.1**, different dibromides, the products analogous to **114.5** and **114.7** are obtained.

Example 115



The preparation of compounds of the invention having phosphonate groups and intermediate compounds useful for their synthesis are illustrated herein. A β -ketonitrile, **115.2**, is generated from a phenylacetic acid, **115.1**, by condensation with a malononitrile ester under Claisen conditions. Reaction with hydroxylamine provides the 5-amino-1,2-oxazole which, upon condensation with cyanomorpholine provides the desired SMP-114 analogs.

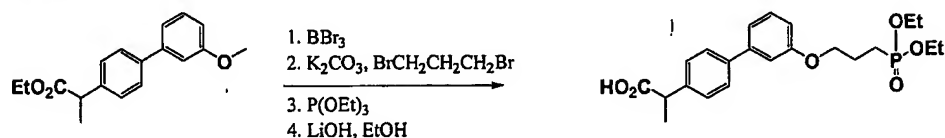
Example 116



The anisole derivative, **116.1**, is demethylated by treatment with a Lewis acid such as boron tribromide. The resulting phenol is alkylated with *E*-1,4-dibromobutene and the resulting monobromide is heated with triethylphosphite in a solvent such as toluene (or other Arbuzov reaction conditions: see Engel, R.,

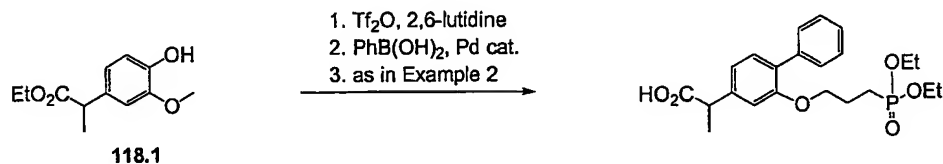
“Synthesis of Carbon-phosphorus Bonds,” CRC press, 1988) to generate the diethyl ester of the desired phosphonic acid, 116.2. Saponification of the carboxylate ester gives the phenylacetic acid ready for incorporation into the synthesis of SMP-114 analogs.

Example 117



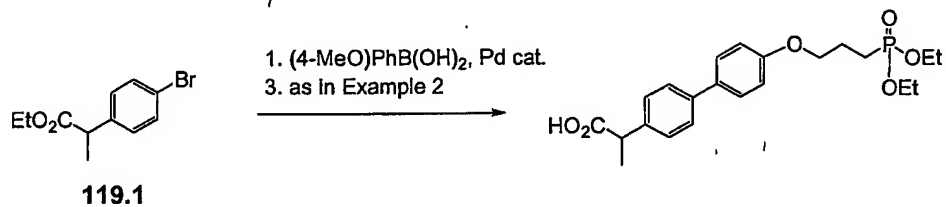
The preparation of compounds of the invention having phosphonate groups can be prepared the procedure of Example 116 and substituting 1,3-dibromopropane in place of the 1,4-dibromobutene.

Example 118

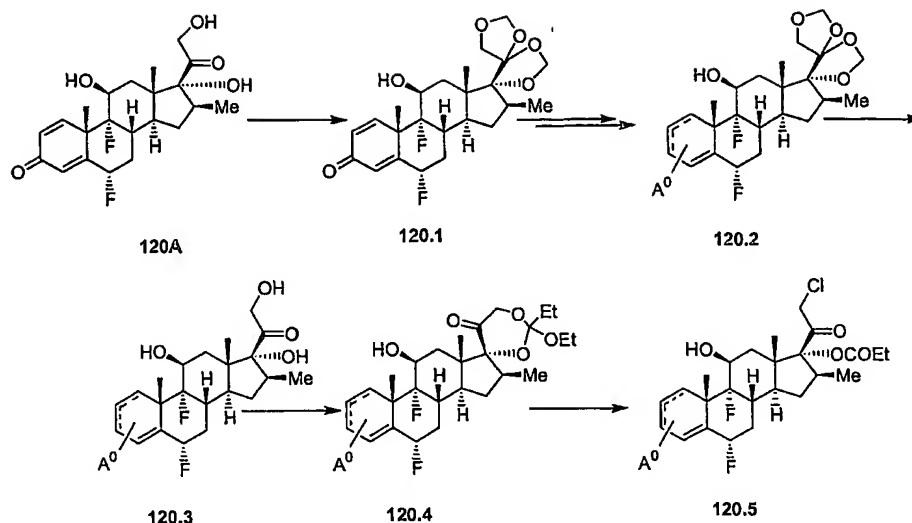


The free phenol in ethyl homovanillate, 118.1, is converted to the aryl triflate, and the biphenyl motif is generated by Suzuki coupling with phenylboronic acid (see *Chem. Rev.*, 1995, 95, 2457). The remaining steps are analogous to those described in Example 116.

Example 119



Ethyl 4-bromophenylacetate, 119.1, is coupled with 4-methoxyphenylboronic acid using the Suzuki method (see above). The remaining steps are analogous to those described in Example 116.

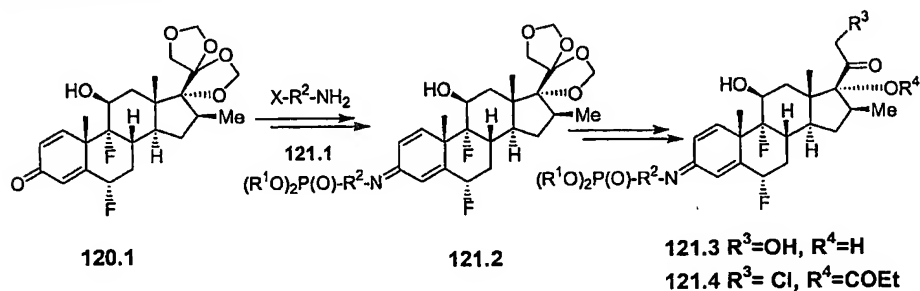
Example 120

6 α ,9 α -Difluoro-16 β -methyl-11 β ,17 α ,21-trihydroxypregn-1,4-dien-3,21-dione, **120A**, (U.S. Patent No. 4,619,921) is reacted with paraformaldehyde and an acid catalyst such as hydrochloric acid, as described in "Protective Groups in Organic Synthesis," by T.W. Greene and P.G.M. Wuts, Wiley, Second Edition 1990, p. 223, to yield the BMD derivative, **120.1**. The phosphonate moiety is then introduced, using the procedures described below, to produce the phosphonate ester **120.2**. The BMD moiety is then hydrolyzed, for example by treatment with 50% aqueous acetic acid, as described in "Protective Groups in Organic Synthesis," by T.W. Greene and P.G.M. Wuts, Wiley, Second Edition 1990, p. 223, to afford the triol, **120.3**. The latter compound is then converted into the 17,21-cyclic orthoester, **120.4**, using the procedure described in *Chem. Pharm. Bull.*, 1986, 34, 1613. The substrate is reacted in dimethylformamide at 70°C with two molar equivalents of triethyl orthopropionate and a catalytic amount of p-toluenesulfonic acid. The product is then reacted with an excess of trimethylsilyl chloride in dimethylformamide at ambient temperature to produce the 21-chloro 17-propionate product, **120.5**.

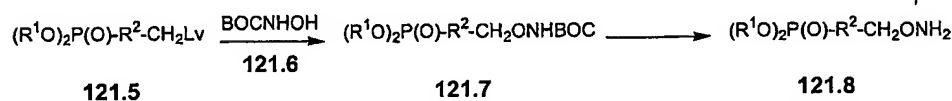
Alternatively, the substrate, **120.3**, is converted into the product, **120.5**, by means of the method described in *J. Med. Chem.*, 1987, 30: 1581. In this procedure, the 21-hydroxy group is activated by conversion to the 21-mesylate, by reaction with mesyl chloride in pyridine; the mesylate group is then displaced to yield the 21-chloro intermediate, by reaction with lithium chloride in dimethylformamide, and the 17-hydroxyl group is esterified to give the 21-

chloro-17-propionate derivative, 120.5. The selective acylation of the 17 α hydroxyl group in the presence of an 11 β hydroxyl group is described in *J. Med. Chem.*, 1987, 30: 1581.

Example 121



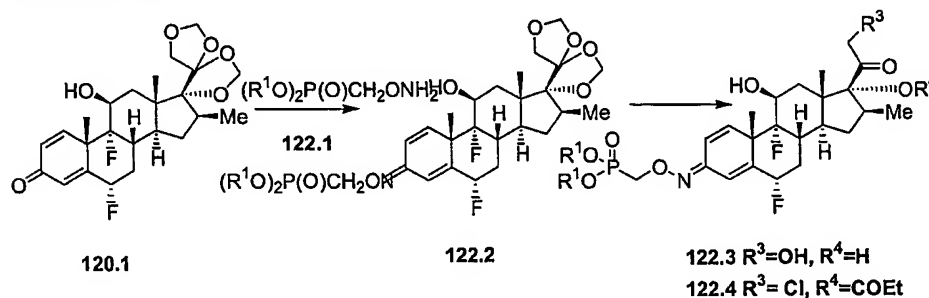
The BMD-protected derivative, 120.1, is reacted with an amine or hydroxylamine, 121.1, in which R² is an alkyl, alkenyl, cycloalkyl or cycloalkenyl group, optionally incorporating a heteroatom O, S or N, or a functional group such as an amide, ester, oxime, sulfoxide or sulfone etc, or an optionally substituted aryl, heteroaryl or aralkyl group, optionally incorporating a heteroatom O, S or N, and X is either a phosphonate group or a group which is subsequently converted into a phosphonate-containing substituent. For example, X is dialkylphosphono, bromo, hydroxy, amino, carboxy and the like. The reaction is conducted between equimolar amounts of the reactants in an aprotic solvent such as pyridine or xylene, or in an alcoholic solvent such as ethanol, optionally in the presence of an acid catalyst, to give the imine or oxime. The preparation of oximes of steroidal 3-ketones is described in *Anal. Bioch.*, 1978, 86, 133 and in *J. Mass. Spectrom.*, 1995, 30, 497. The BMD-protected side-chain compound, 121.2, is then converted into the triol, 121.3, and then to the 21-chloro 17 propionate product, 121.4, as described above.



The preparation of hydroxylamine ethers incorporating a phosphonate group is illustrated above. A phosphonate, 121.5, in which Lv is a leaving group such as bromo or trifluoromethylsulfonyloxy, is reacted with BOC-hydroxylamine, 121.6, (Aldrich) to produce the ether, 121.7. The reaction is

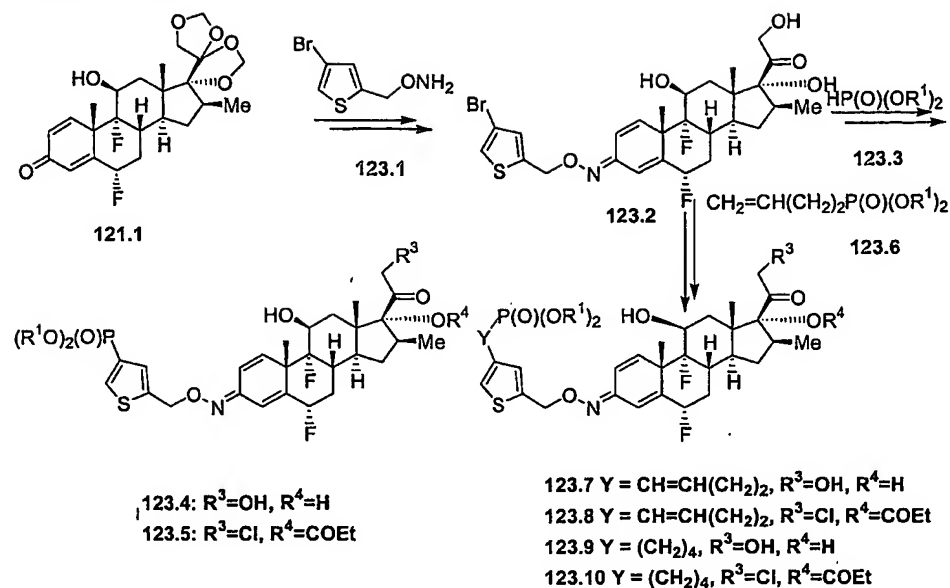
conducted between equimolar amounts of the reactants in a polar solvent such as dimethylformamide or tetrahydrofuran, in the presence of a base such as potassium hydroxide or dimethylaminopyridine. Deprotection, for example by treatment with trifluoroacetic acid, then gives the hydroxylamine ether, 121.8.

Example 122



The substrate, 120.1, is reacted with a dialkyl phosphonomethyl hydroxylamine, 122.1, prepared as described above from a dialkyl trifluoroethylsulfonyloxymethyl phosphonate (*Tet. Lett.*, 1986, 27:1477) and BOC-hydroxylamine, to afford the oxime, 122.2. Deprotection then affords the triol, 122.3, from which the 21-chloro 17-propionate compound, 122.4, is prepared. The oxime forming reaction is performed at ambient temperature in ethanol-acetic acid solution between equimolar amounts of the reactants.

Using the above procedures, but employing, in place of the hydroxylamine ether, 122.1, different oxime ethers, 121.1, the corresponding products, 121.4 are obtained.

Example 123

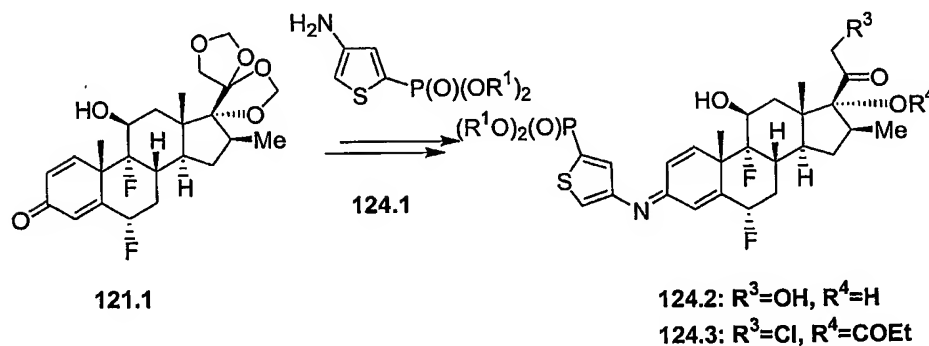
The dienone, 121.1, is reacted, as described above, with O-(4-bromo-2-thienylmethoxy)hydroxylamine, 123.1, prepared as described above from 4-bromo-2-bromomethylthiophene (WO 94/20456) and BOC-protected hydroxylamine, to give, after deprotection of the side-chain, the oxime, 123.2. The product is then reacted, in the presence of a palladium catalyst, with a dialkyl phosphite, 123.3, to afford the phosphonate, 123.4. The preparation of arylphosphonates by means of a coupling reaction between aryl bromides and dialkyl phosphites is described in *J. Med. Chem.* 1992, 35, 1371. The reaction is performed in an inert solvent such as toluene, in the presence of a base such as triethylamine and a catalytic amount of tetrakis(triphenylphosphine)-palladium(0). The 21-hydroxy compound, 123.4, is then converted, as described herein, into the 21-chloro 17-propionate derivative, 123.5.

Alternatively, the bromo compound, 123.2, is coupled with a dialkyl butenyl phosphonate, 123.6, (*Org. Lett.*, 2001, 3, 217) to afford the phosphonate, 123.7. The coupling of aryl halides with olefins by means of the Heck reaction is described, for example, in F. A. Carey and R. J. Sundberg, *Advanced Organic Chemistry*, 503ff (Plenum, 2001) and in *Acc. Chem. Res.*, 1979, 12, 146. The aryl bromide and the olefin are coupled in a polar solvent such as dimethylformamide or dioxan, in the presence of a palladium(0) catalyst such as tetrakis(triphenylphosphine)palladium(0) or palladium(II) catalyst such as

palladium(II) acetate, and optionally in the presence of a base such as triethylamine or potassium carbonate. Optionally, the double bond present in the product, **123.7**, is reduced, for example by reaction with diimide, to produce the saturated analog, **123.9**. The reduction of olefinic bonds is described in R. C. Larock, *Comprehensive Organic Transformations*, 6ff (VCH 1989). The transformation is effected by means of catalytic hydrogenation, for example using a palladium on carbon catalyst and hydrogen or a hydrogen donor, or by the use of diimide or diborane. The products, **123.7** and **123.9**, are then converted into the 21-chloro 17-propionate analogs, **123.8** and **123.10**.

Using the above procedures, but employing, in place of the bromothiénylmethoxy reagent, **123.1**, different bromo-substituted aryl or heteroaryl alkoxy hydroxylamines, and/or different dialkyl alkenyl phosphonates, the products analogous to the compounds, **123.5**, **123.8** and **123.10** are obtained.

Example 124

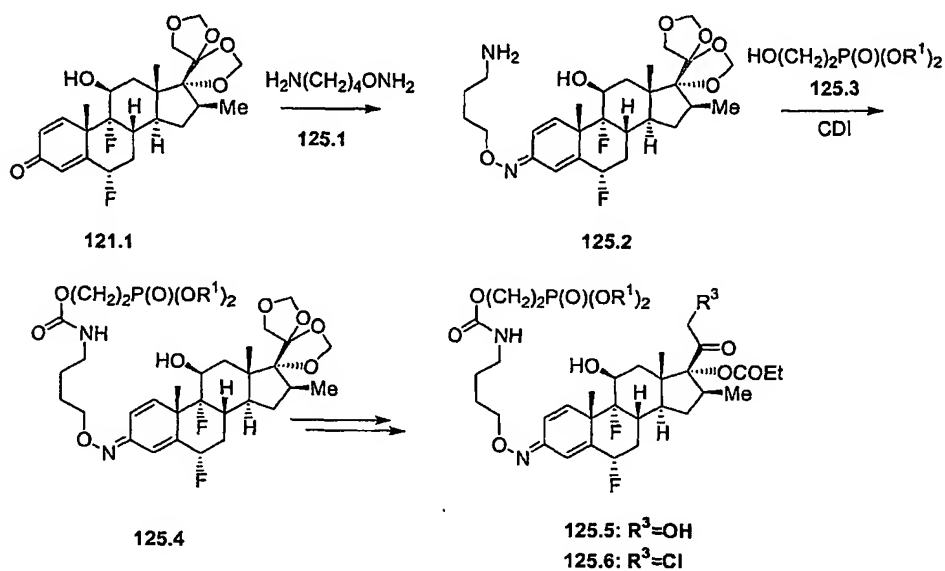


The substrate, **121.1**, is reacted with a dialkyl 4-amino-2-thienyl phosphonate, **124.1**, prepared by the palladium-catalyzed coupling, as described above, between 4-amino-2-bromothiophene (*Tet.*, **1987**, *43*, 3295) and a dialkyl phosphite, to give, after deprotection, the imine product, **124.2**. The imine forming reaction is conducted in a hydrocarbon solvent such as toluene or xylene, at reflux temperature, in the presence of a basic catalyst such as sodium methoxide, or an acid catalyst such as p-toluenesulfonic acid, under azeotropic

conditions. The product is then converted into the 21-chloro 17-propionate compound, 124.3.

Using the above procedures, but employing, in place of the 4-aminothienyl phosphonate, 124.1 different amino-substituted aryl or heteroaryl phosphonates, products analogous to 124.3 are obtained.

Example 125

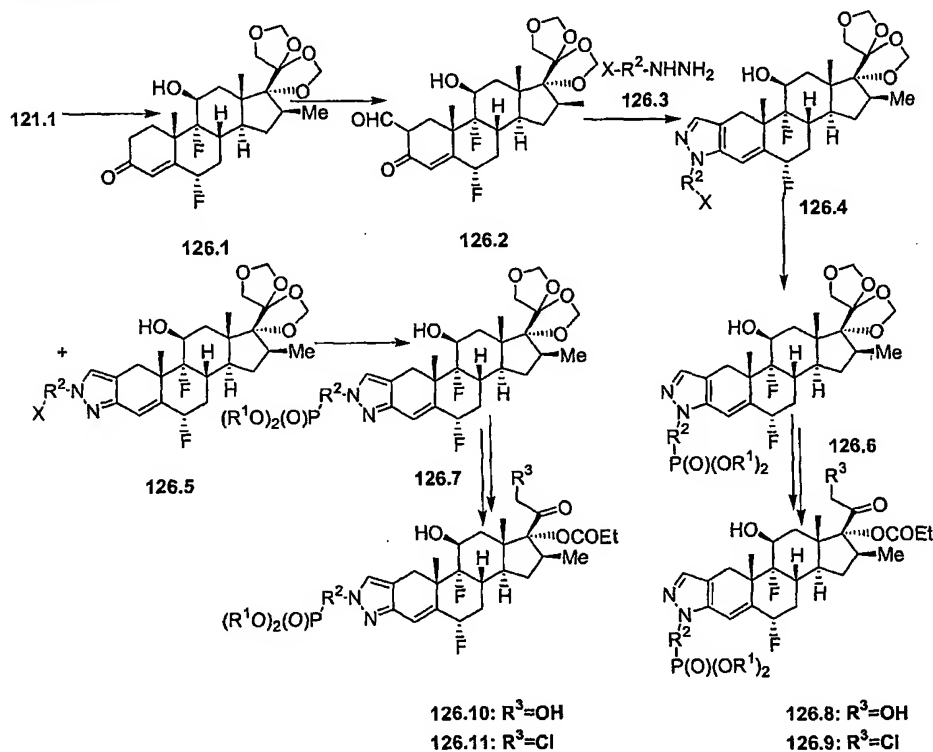


The dienone, 121.1, is reacted with O-(4-aminobutyl)hydroxylamine, 125.1, (*Pol. J. Chem.*, 1981, 55, 1163) to yield the oxime, 125.2. The reaction of steroidal 1,4-dien-3-ones with substituted hydroxylamines is described in *J. Steroid Bioch.*, 1976, 7, 795; the reaction is performed between equimolar amounts of the reactants in a polar organic solvent such as pyridine or methanol, optionally in the presence of acetic acid or sodium acetate. The product is then coupled with a dialkyl 2-hydroxyethyl phosphonate, 125.3, (Epsilon) and carbonyl diimidazole, to yield the carbamate oxime, 125.4. The preparation of carbamates is described in A. R. Katritzky, *Comprehensive Organic Functional Group Transformations*, 6, 416ff (Pergamon, 1995), and in S. R. Sandler and W. Karo, *Organic Functional Group Preparations*, 260ff (Academic Press, 1986). In the procedure, the amine is reacted in an inert aprotic solvent such as dichloromethane or tetrahydrofuran, with phosgene or a functional equivalent thereof, such as carbonyl diimidazole, triphosgene, pentafluorophenyl carbonate

and the like, to afford the corresponding activated acylamine. The latter compound is then reacted with an alcohol to yield the carbamate. The carbamate product, **125.4**, is then converted, as described herein, into the 21-chloro 17-propionate product, **125.6**.

Using the above procedures, but employing, in place of the hydroxylamine, **125.3**, different amino-substituted hydroxylamines, and/or different hydroxy-substituted phosphonates, the products analogous to **125.6** are obtained.

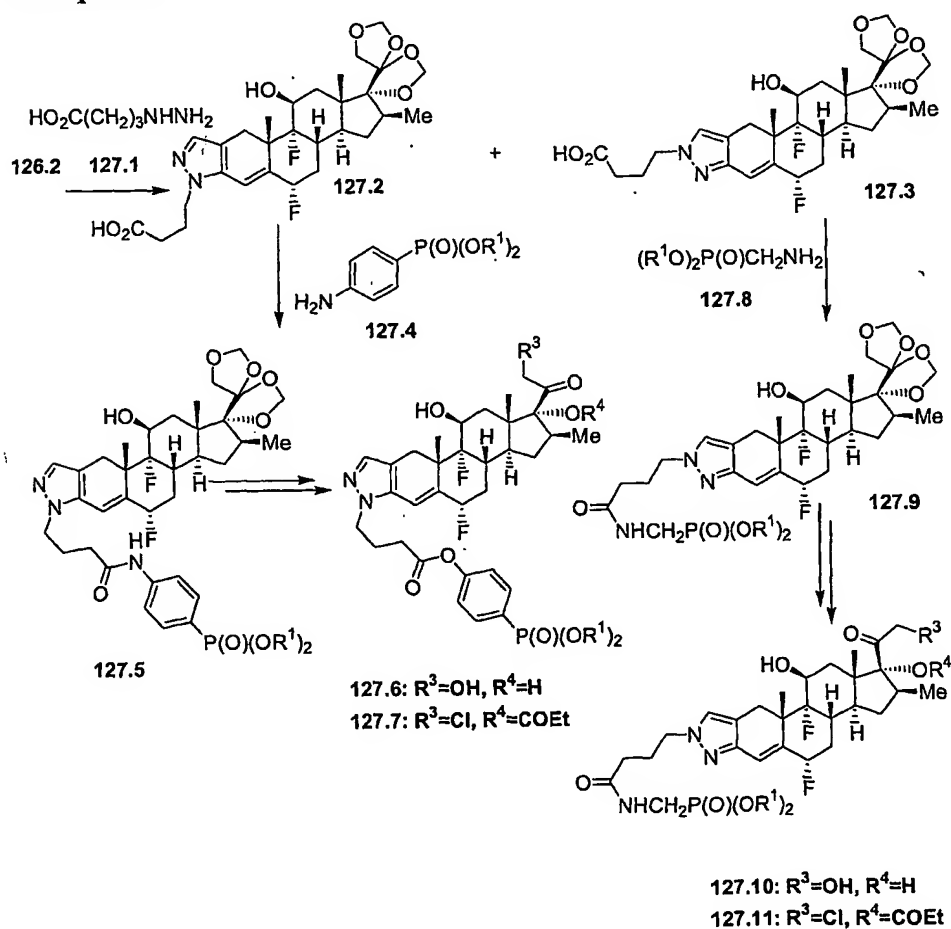
Example 126



The BMD-protected dienone, **121.1**, is reduced to afford the 1,2-dihydro product, **126.1**. The catalytic hydrogenation reaction is effected by the use of tris(triphenylphosphine)rhodium (I) chloride, for example as described in *J. Med. Chem.*, **2001**, *44*, 602. The product is then reacted with ethyl formate and a base such as sodium hydride, in an inert solvent such as toluene or dimethylformamide, as described in *J. Am. Chem. Soc.*, **1964**, *86*, 1520, to afford the 2-formyl product, **126.2**. This compound is then reacted with an alkyl,

aralkyl, aryl or heteroaryl hydrazine, **126.3**, in which the substituent X is either a phosphonate group or a group which is subsequently transformed into a phosphonate-containing substituent. For example, X is dialkylphosphono, bromo, hydroxy, amino, carboxyl and the like. The reaction yields the isomeric 2'- and 1'-aryl pyrazoles, **126.4** and **126.5**. The pyrazole-forming reaction is performed between equimolar amounts of the reactants in an acidic solvent such as acetic acid, as described in *J. Am. Chem. Soc.*, 1964, 86, 1520. The pyrazoles, **126.4** and **126.5**, are then transformed, for example by the procedures described in Examples 122 and 123, via the BMD-protected intermediates, **126.6** and **126.7**, into the 21-chloro 17-propionate phosphonates, **126.9** and **126.11**.

Example 127



The ketoaldehyde, **126.2**, is reacted, as described above, with 3-carboxypropyl hydrazine, **127.1**, (*Ind. J. Exp. Biol.*, 1994, 32, 218) to give the

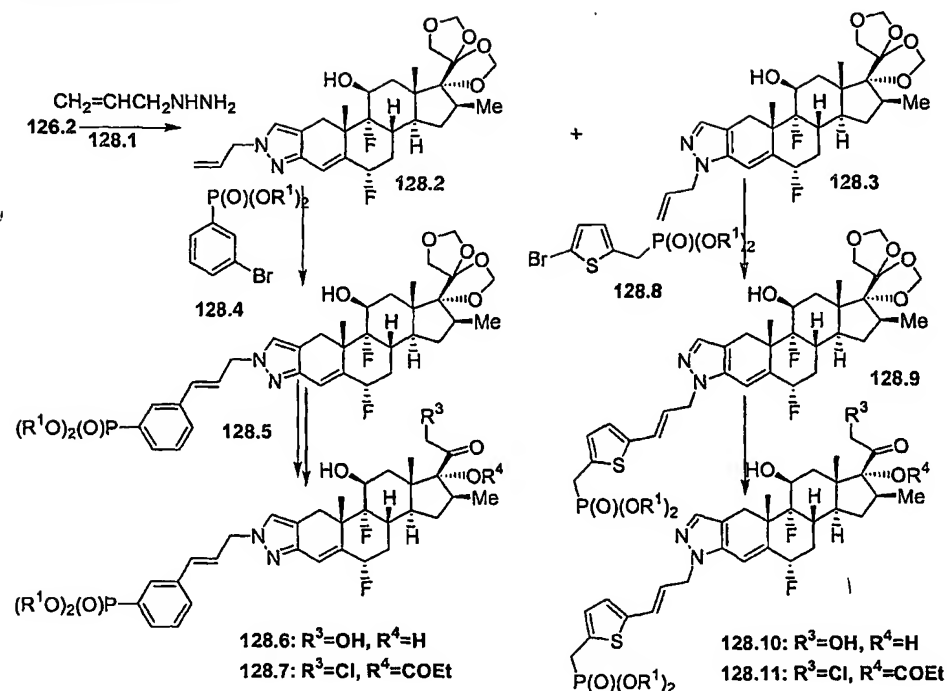
pyrazoles, 127.2 and 127.3. The 2'-substituted isomer, 127.2, is then reacted in dimethylformamide solution at ambient temperature with one molar equivalent of a dialkyl 4-aminophenyl phosphonate, 127.4, (Epsilon) and dicyclohexyl carbodiimide, to yield the amide, 127.5. The preparation of amides from carboxylic acids and derivatives is described, for example, in S.R. Sandler and W. Karo, *Organic Functional Group Preparations*, 274 (Academic Press, 1968), and R. C. Larock, *Comprehensive Organic Transformations*, 972ff (VCH, 1989). The carboxylic acid is reacted with the amine in the presence of an activating agent, such as, for example, dicyclohexylcarbodiimide or diisopropylcarbodiimide, optionally in the presence of, for example, hydroxybenztriazole, N-hydroxysuccinimide or N-hydroxypyridone, in a non-protic solvent such as, for example, pyridine, DMF or dichloromethane, to afford the amide.

Alternatively, the carboxylic acid may first be converted into an activated derivative such as the acid chloride, anhydride, mixed anhydride, imidazolidine and the like, and then reacted with the amine, in the presence of an organic base such as, for example, pyridine, to afford the amide.

The conversion of a carboxylic acid into the corresponding acid chloride can be effected by treatment of the carboxylic acid with a reagent such as, for example, thionyl chloride or oxalyl chloride in an inert organic solvent such as dichloromethane, optionally in the presence of a catalytic amount of dimethylformamide. The BMD protecting group is then removed and the product is converted into the 21-chloro 17-propionate product, 127.7.

The 1'-substituted pyrazole, 127.3, is coupled, as described above, with a dialkyl aminomethyl phosphonate, 127.8 (Interchim), to afford the amide, 127.9. The product, 127.9, is then deprotected to give the triol, 127.10, and the latter compound is transformed into the 21-chloro 17-propionate, 127.11.

Using the above procedures, but employing different amino-substituted phosphonates, and/or different carboxy-substituted hydrazines, the products analogous to 127.7 and 127.11 are obtained. The functionalization procedures are interchangeable between the pyrazole substrates, 127.2 and 127.3.

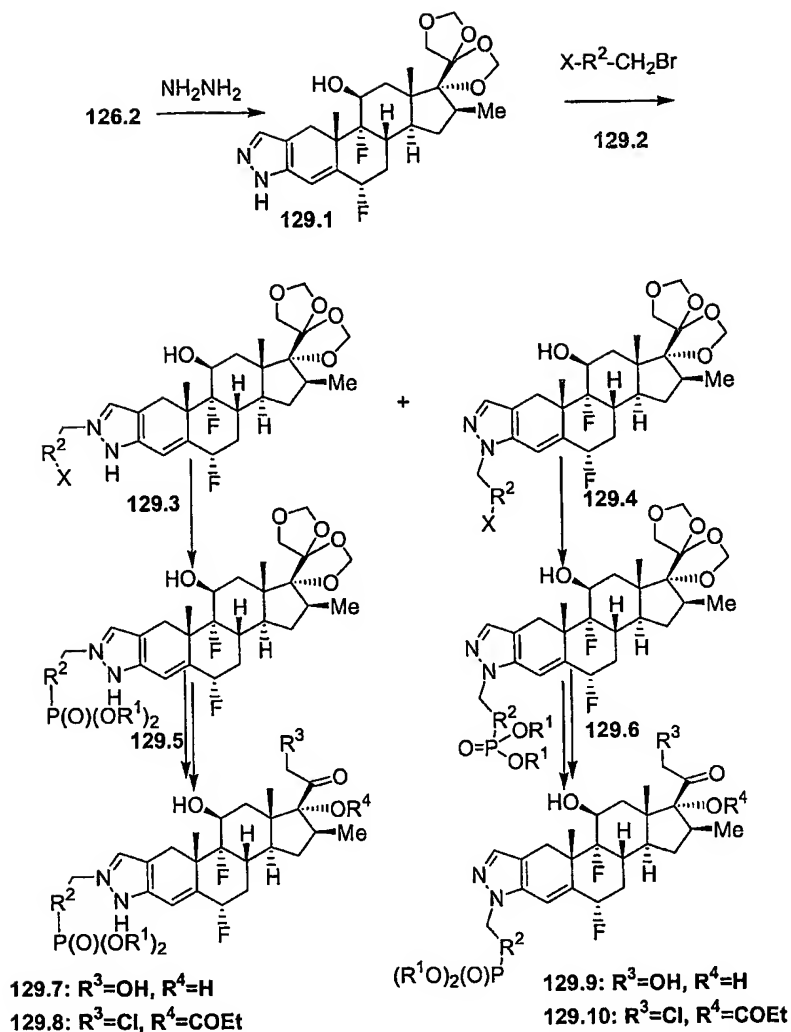
Example 128

The ketoaldehyde, 126.2, is reacted, as described above, with allylhydrazine, 128.1, (*Zh. Org. Khim.*, 1967, 3, 983) to produce the pyrazoles, 128.2 and 128.3. The 1'-substituted isomer, 128.2, is coupled, as described herein, with a dialkyl 3-bromophenyl phosphonate, 128.4, (Epsilon) to give the phosphonate, 128.5. The product is then deprotected to afford the triol, 128.6, which is converted into the 21-chloro 17-propionate compound, 128.7.

The 2'-substituted pyrazole, 128.3, is coupled, as described above, with a dialkyl 5-bromo-2-thienyl phosphonate, 128.8, (*Syn.*, 2003, 455) to prepare the phosphonate, 128.9, which is deprotected, and the product is converted into the 21-chloro 17-propionate analog, 128.11.

Using the above procedures, but employing, in place of the propenylhydrazine, 128.1, different alkenyl hydrazines, and/or different dialkyl bromo-substituted phosphonates, the products analogous to the compounds, 128.7 and 128.11 are obtained.

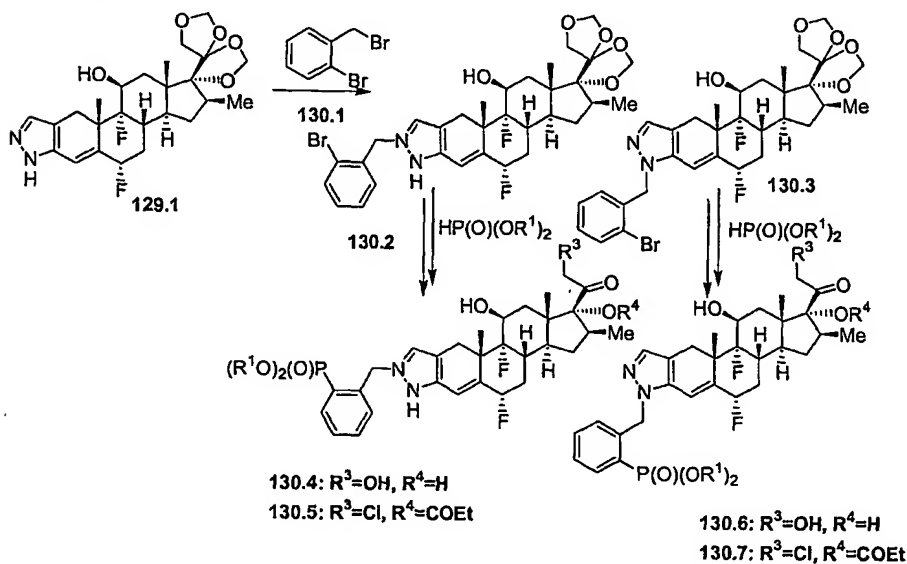
Example 129



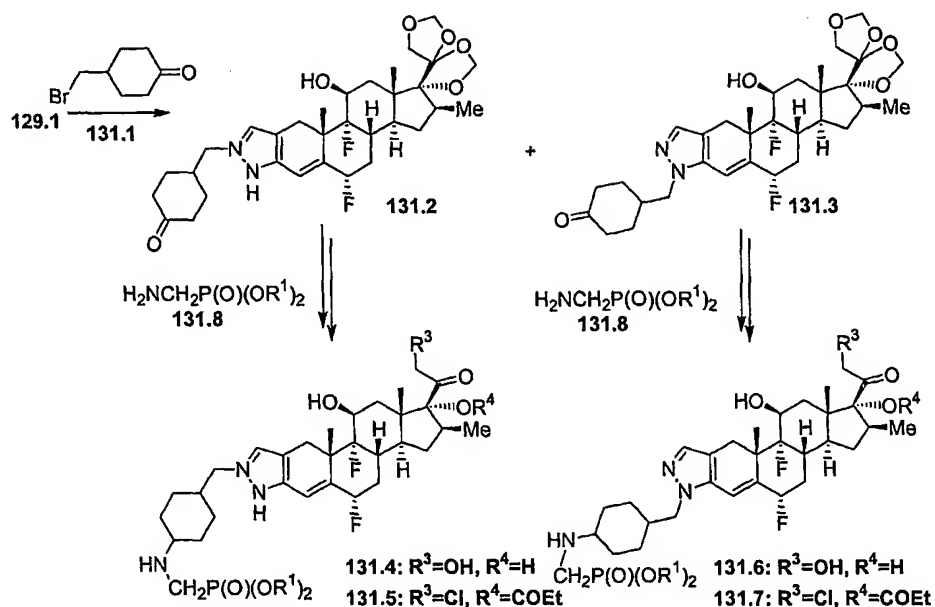
The ketoaldehyde, **126.2**, is reacted with hydrazine, to afford the pyrazole derivative, **129.1**. The reaction of steroidal 2-formyl-3-ketones with hydrazine is described in *J. Am. Chem. Soc.*, 1964, 86, 1520. The reaction is performed in acetic acid at ambient temperature. The pyrazole product is then reacted with a bromomethyl compound, **129.2**, in which R^2 and X are as defined above, to yield the alkylation products, **129.3** and **129.4**. The alkylation of substituted pyrazoles is described, for example, in T. L. Gilchrist, *Heterocyclic Chemistry*, 309 (Longman, 1992). The reaction is performed between equimolar amounts of the substrates in a polar solvent such as dimethylformamide or tetrahydrofuran, in the presence of a base such as dimethylaminopyridine, lithium hexamethyldisilazide and the like. The products, **129.3** and **129.4**, are,

except in cases where X is dialkylphosphono, converted into the phosphonates, 129.5 and 129.6, using the procedures described herein, and deprotection/chlorination/acylation then affords the 21-chloro 17-propionate compounds, 129.8 and 129.10.

Example 130



Following the procedure described in Example 129, the pyrazole, 129.1, is reacted with 2-bromobenzyl bromide, 130.1, to give the pyrazoles, 130.2 and 130.3. The products are then coupled, as described above, with a dialkyl phosphite, to afford, after side-chain deprotection and modification, as described herein, the 21-chloro 17 propionates, 130.5 and 130.7.

Example 131

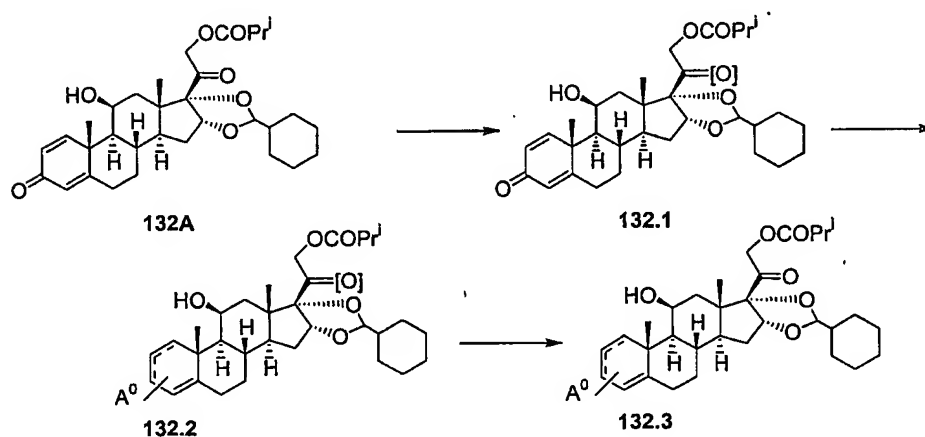
Following the procedure described in Example 129, the pyrazole, 129.1, is reacted in tetrahydrofuran solution with 4-bromomethyl cyclohexanone, 131.1, (WO 97/37959) to give the alkylation products, 131.2 and 131.3. The 1'-substituted isomer, 131.2, is then reacted, in a reductive amination reaction, with a dialkyl aminomethyl phosphonate, 131.8, (Interchim) and sodium cyanoborohydride, to yield, after deprotection and side-chain modification, the 21-chloro 17-propionate, 131.5.

The preparation of amines by means of reductive amination procedures is described, for example, in R. C. Larock, *Comprehensive Organic Transformations*, 421 (VCH, 1989), and in F.A. Carey and R. J. Sundberg, *Advanced Organic Chemistry*, Part B, 269 (Plenum, 2001). In this procedure, the amine component and the aldehyde or ketone component are reacted together in the presence of a reducing agent such as, for example, borane, sodium cyanoborohydride, sodium triacetoxyborohydride or diisobutylaluminum hydride, optionally in the presence of a Lewis acid, such as titanium tetraisopropoxide, as described in *J. Org. Chem.*, 1990, 55, 2552.

The 2'-substituted pyrazole, 131.3, is subjected to the same series of reaction to give the amine phosphonate, 131.7.

Using the above procedures, but employing different bromomethyl-substituted aldehydes or ketones, and/or different amino-substituted phosphonates, the products analogous to 131.5 and 131.7 are obtained.

Example 132



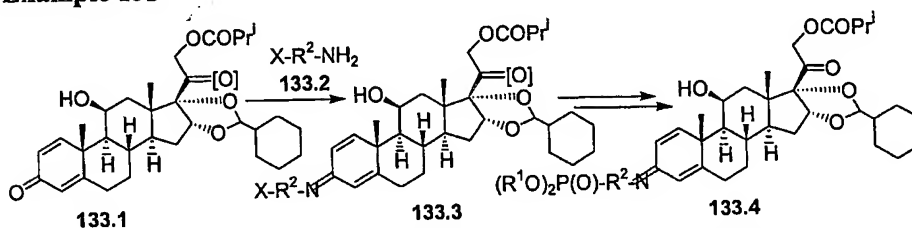
A protection-deprotection sequence in which the 20-ketone group of Ciclesonide, 132A, (U.S. Patent No. 5,482,934) is protected to afford the derivative, 132.1. The ketone is protected, for example, by conversion to the cyclic ethylene ketal, by reaction in toluene solution at reflux temperature with ethylene glycol and an acid catalyst, as described in *J. Am. Chem. Soc.*, 1955, 77, 1904. Deprotection is effected by reaction with pyridinium tosylate in aqueous acetone, as described in *J. Chem. Soc. Chem. Comm.*, 1987, 1351.

Alternatively, the 20-ketone is protected by conversion to the N, N-dimethylhydrazone. The dimethyl hydrazone is prepared by the reaction of the ketone, 132A, with N,N-dimethylhydrazine in ethanol-acetic acid, as described in *Org. Syn.*, 1970, 50, 102. The group is removed by treatment with sodium acetate and acetic acid in aqueous tetrahydrofuran, as described in *J. Am. Chem. Soc.*, 1979, 101, 5841.

Alternatively, the 20-ketone is protected as the diethylamine adduct. In this procedure, the substrate, 132A, is reacted with titanium tetrakis(diethylamide), as described in *J. Chem. Soc. Chem. Comm.*, 1983, 406, to afford the adduct. The ketone is deprotected by reaction with water in an aqueous organic solvent.

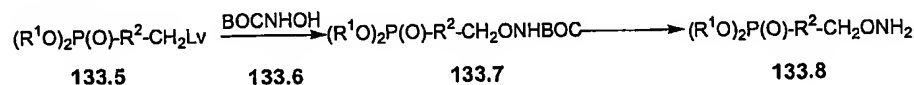
The protected compound, 132.1, is then converted into the phosphonate-containing analog, 132.2, using the procedures described below, and the protecting group is then removed, as described above, to give the phosphonate, 132.3.

Example 133



The protected derivative, 133.1, is reacted with an amine or hydroxylamine, 133.2, in which R^2 is an alkyl, alkenyl, cycloalkyl or cycloalkenyl group, optionally incorporating a heteroatom O, S or N, or a functional group such as an amide, ester, oxime, sulfoxide or sulfone etc, or an optionally substituted aryl, heteroaryl or aralkyl group, optionally incorporating a heteroatom O, S or N, and X is either a phosphonate group or a group which is subsequently converted into a phosphonate-containing substituent. For example, X is dialkylphosphono, bromo, hydroxy, amino, carboxy and the like. The reaction is conducted between equimolar amounts of the reactants in an aprotic solvent such as pyridine or xylene, or in an alcoholic solvent such as ethanol, optionally in the presence of an acid catalyst, to give the imine or oxime, 133.3. The preparation of oximes of steroidal 3-ketones is described in *Anal. Bioch.*, 1978, 86, 133 and in *J. Mass. Spectrom.*, 1995, 30, 497. The protecting group is then removed, as described herein, to afford the 20-keto phosphonate product, 133.4.

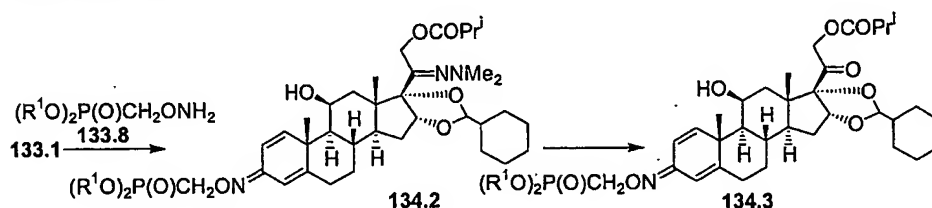
Example 133A



The preparation of hydroxylamine ethers incorporating a phosphonate group is illustrated herein. In this procedure, a phosphonate, 133.5, in which Lv is a leaving group such as bromo or trifluoromethylsulfonyloxy, is reacted with BOC-hydroxylamine, 133.6, (Aldrich) to produce the ether, 133.7. The reaction

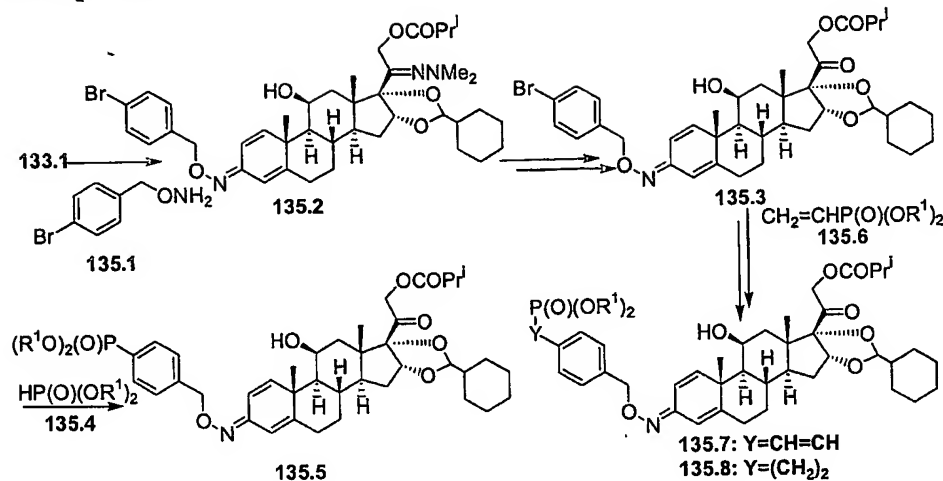
is conducted between equimolar amounts of the reactants in a polar solvent such as dimethylformamide or tetrahydrofuran, in the presence of a base such as potassium hydroxide or dimethylaminopyridine. Deprotection, for example by treatment with trifluoroacetic acid, then gives the hydroxylamine ether, **133.8**. The above procedure is also employed for the preparation of substituted hydroxylamines which are precursors to phosphonates.

Example 134



The substrate, **133.1**, in which the 20-ketone is protected as the dimethyl hydrazone derivative, is reacted with a dialkyl phosphonomethyl hydroxylamine, **133.8**, prepared as described above from a dialkyl trifluoromethylsulfonyloxymethyl phosphonate (*Tetrahedron Lett.*, **1986**, 27, 1477) and BOC-hydroxylamine, to afford the oxime, **134.2**. Deprotection, as described in Example 132, then affords the 20-keto phosphonate, **134.3**. The oxime forming reaction is performed at ambient temperature in ethanol-acetic acid solution between equimolar amounts of the reactants.

Using the above procedures, but employing, in place of the hydroxylamine ether, **133.8**, different oxime ethers, **133.2**, the corresponding products, **133.4** are obtained.

Example 135

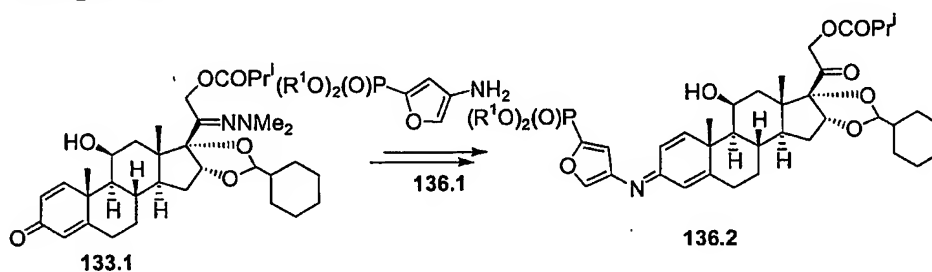
The dienone, 133.1, in which the 20-ketone is protected as the dimethyl hydrazone, is reacted, as described above, with O-(4-bromobenzyloxy)-hydroxylamine, 135.1, prepared as described above from 4-bromobenzyl bromide and BOC-protected hydroxylamine, 133.6, to give the oxime, 135.2. The protecting group is then removed to yield the 20-keto product, 135.3. The latter product is then reacted, in the presence of a palladium catalyst, with a dialkyl phosphate, 135.4, to afford the phosphonate, 135.5. The preparation of arylphosphonates by means of a coupling reaction between aryl bromides and dialkyl phosphites is described in *J. Med. Chem.*, **1992**, *35*, 1371. The reaction is performed at ca. 100°C in an inert solvent such as toluene, in the presence of a base such as triethylamine and a catalytic amount of tetrakis(triphenylphosphine)palladium(0).

Alternatively, the bromo compound, 135.3, is coupled with a dialkyl vinyl phosphonate, 135.6, (Aldrich) to afford the phosphonate, 135.7. The coupling of aryl halides with olefins by means of the Heck reaction is described, for example, in F. A. Carey and R. J. Sundberg, *Advanced Organic Chemistry* 503ff (Plenum, 2001) and in *Acc. Chem. Res.*, **1979**, *12*, 146. The aryl bromide and the olefin are coupled in a polar solvent such as dimethylformamide or dioxan, in the presence of a palladium(0) catalyst such as tetrakis(triphenylphosphine)palladium(0) or palladium(II) catalyst such as palladium(II) acetate, and optionally in the presence of a base such as triethylamine or potassium carbonate. Optionally, the styrenoid double bond present in the product, 135.7,

is reduced, for example by reaction with diimide, to produce the saturated analog, 135.8. The reduction of olefinic bonds is described in R. C. Larock, *Comprehensive Organic Transformations* 6ff (VCH 1989). The transformation is effected by means of catalytic hydrogenation, for example using a palladium on carbon catalyst and hydrogen or a hydrogen donor, or by the use of diimide or diborane.

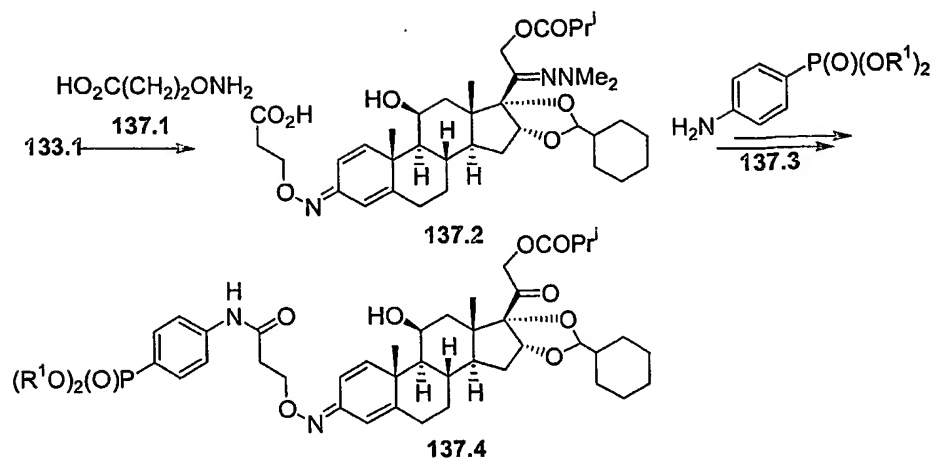
Using the above procedures, but employing, in place of the bromobenzyl-oxy reagent, 135.1, different bromo-substituted aryl or heteroaryl alkoxy hydroxylamines, and/or different dialkyl alkenyl phosphonates, products analogous to the compounds, 135.5, 135.7 and 135.8 are obtained.

Example 136



The substrate, 133.1, in which the 20-ketone is protected as the dimethylhydrazone, is reacted with a dialkyl 4-amino-2-furyl phosphonate, 136.1, prepared by the palladium catalyzed coupling reaction, as described above, between 4-amino-2-bromofuran (*Tetrahedron Lett.*, 1987, 43, 3295) and a dialkyl phosphite, to give, after deprotection, the imine product, 136.2. The imine forming reaction is conducted in a hydrocarbon solvent such as toluene or xylene, at reflux temperature, in the presence of a basic catalyst such as sodium methoxide, or an acid catalyst such as p-toluenesulfonic acid, under azeotropic conditions.

Using the above procedures, but employing, in place of the 4-amino-2-furyl phosphonate, 136.1, different amino-substituted aryl or heteroaryl phosphonates, products analogous to 136.2 are obtained.

Example 137

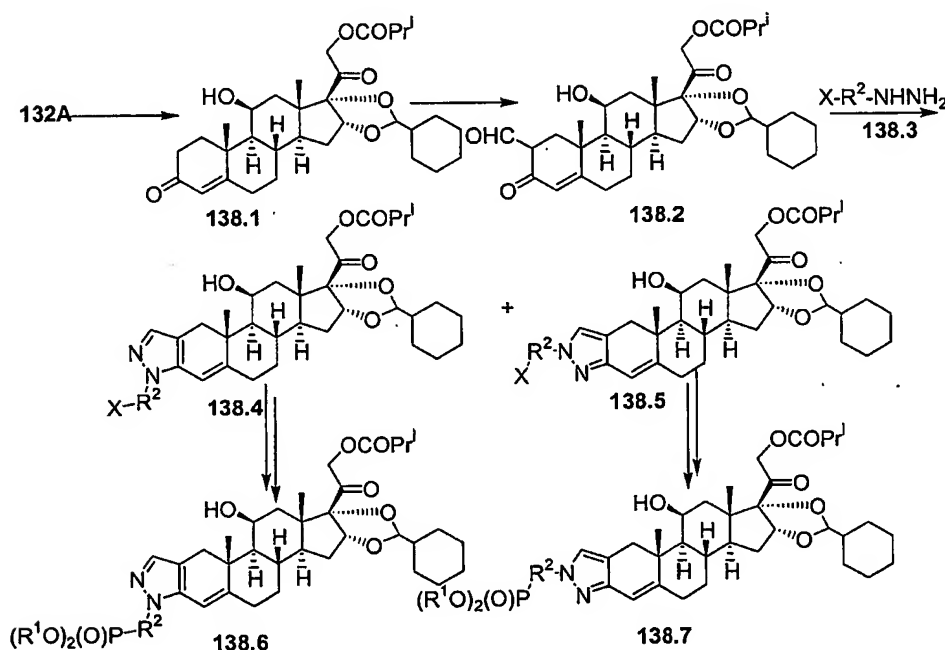
The dienone, 133.1, in which the 20-ketone is protected as the dimethylhydrazone, is reacted with O-(2-carboxyethyl)hydroxylamine, 137.1, (*J. Med. Chem.*, 1990, 33, 1423) to yield the oxime, 137.2. The reaction of steroidal 1,4-dien-3-ones with substituted hydroxylamines is described in *J. Steroid Bioch.*, 1976, 7, 795; the reaction is performed between equimolar amounts of the reactants in a polar organic solvent such as pyridine or methanol, optionally in the presence of acetic acid or sodium acetate. The product, 137.2, is then coupled with a dialkyl 4-aminophenyl phosphonate, 137.3, (Epsilon) and dicyclohexylcarbodiimide, to yield, after deprotection the amide oxime, 137.4. The preparation of amides from carboxylic acids and derivatives is described, for example, in S.R. Sandler and W. Karo, *Organic Functional Group Preparations* 274 (Academic Press, 1968) and R. C. Larock, *Comprehensive Organic Transformations* 972ff (VCH, 1989). The carboxylic acid is reacted with the amine in the presence of an activating agent, such as, for example, dicyclohexylcarbodiimide or diisopropylcarbodiimide, optionally in the presence of, for example, hydroxybenztriazole, N-hydroxysuccinimide or N-hydroxypyridone, in a non-protic solvent such as, for example, pyridine, DMF or dichloromethane, to afford the amide.

Alternatively, the carboxylic acid may first be converted into an activated derivative such as the acid chloride, anhydride, mixed anhydride, imidazolidine and the like, and then reacted with the amine, in the presence of an organic base such as, for example, pyridine, to afford the amide.

The conversion of a carboxylic acid into the corresponding acid chloride can be effected by treatment of the carboxylic acid with a reagent such as, for example, thionyl chloride or oxalyl chloride in an inert organic solvent such as dichloromethane, optionally in the presence of a catalytic amount of dimethylformamide.

Using the above procedures, but employing, in place of the carboxy-substituted hydroxylamine, **137.1**, different carboxy-substituted hydroxylamines, and/or different amino-substituted phosphonates, products analogous to **137.4** are obtained.

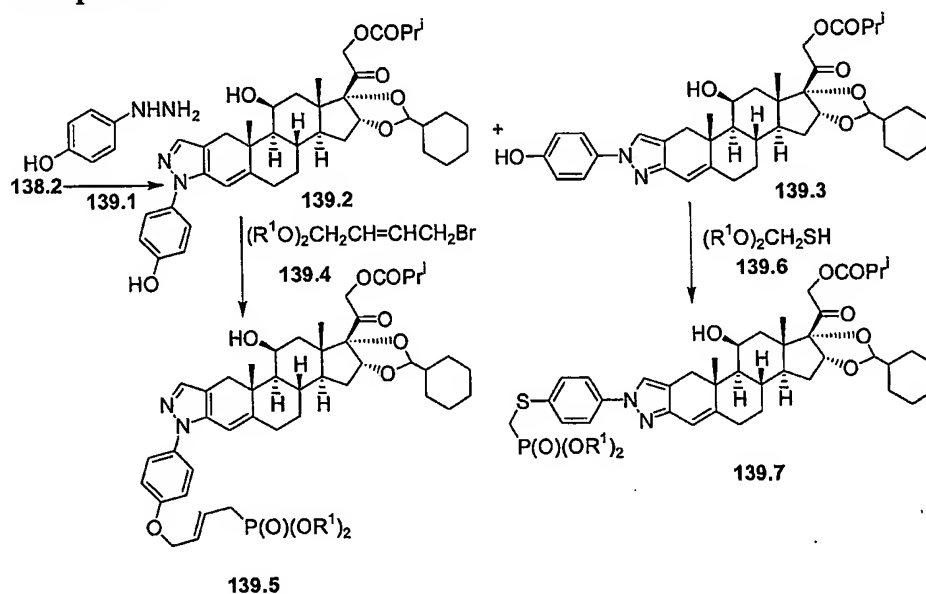
Example 138



The dienone, **132A**, is reduced to afford the 1,2-dihydro product, **138.1**. The catalytic hydrogenation reaction is effected by the use of tris(triphenylphosphine)rhodium (I) chloride, for example as described in *J. Med. Chem.*, **2001**, *44*, 602. The product is then reacted with ethyl formate and a base such as sodium hydride, in an inert solvent such as toluene or dimethylformamide, as described in *J. Am. Chem. Soc.*, **1964**, *86*, 1520, to afford the 2-formyl product, **138.2**. This compound is then reacted with an alkyl, aralkyl, aryl or heteroaryl

hydrazine, 138.3, in which the substituent X is either a phosphonate group or a group which is subsequently transformed into a phosphonate-containing substituent. For example, X is dialkylphosphono, bromo, hydroxy, amino, carboxyl and the like. The reaction yields the isomeric 2'- and 1'-aryl pyrazoles, 138.4 and 138.5. The pyrazole-forming reaction is performed between equimolar amounts of the reactants in an acidic solvent such as acetic acid, as described in *J. Am. Chem. Soc.*, 1964, 86, 1520. The pyrazoles, 138.4 and 138.5, are then transformed, for example by the procedures described in Examples 139 and 140, into the phosphonates, 138.6 and 138.7.

Example 139



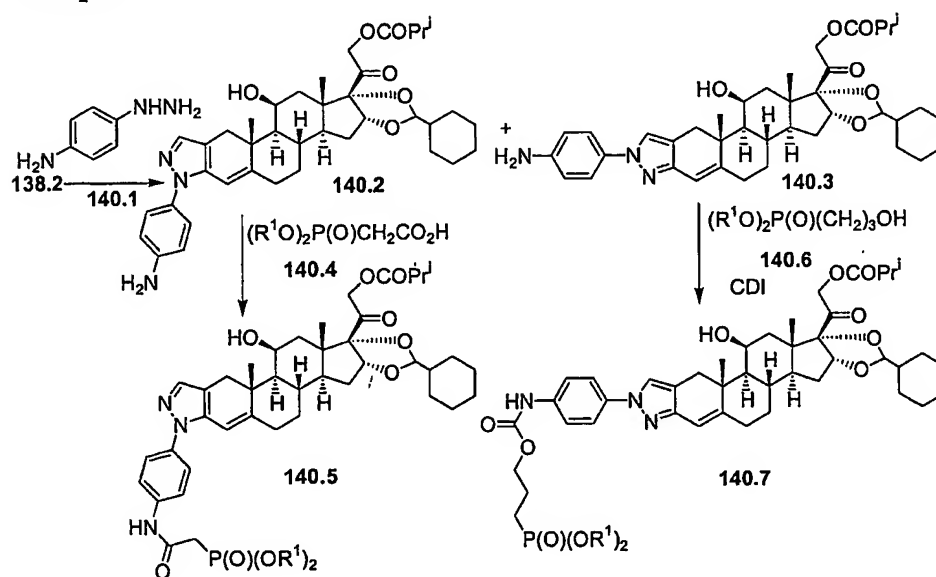
The ketoaldehyde, 138.2, is reacted, as described above, with 4-hydroxyphenyl hydrazine, 139.1, (EP 437105) to give the pyrazoles, 139.2 and 139.3. The 2'-substituted isomer, 139.2, is then reacted in dimethylformamide solution at ca. 70°C with a dialkyl bromobutenyl phosphonate, 139.4, (*J. Med. Chem.*, 1992, 35, 1371) and potassium carbonate, to yield the ether phosphonate, 139.5.

The isomeric pyrazole, 139.3, is reacted, in a Mitsunobu reaction, with a dialkyl mercaptomethyl phosphonate, 139.6, (*J. Med. Chem.*, 1985, 26, 1688) to yield the thioether phosphonate, 139.7. The preparation of aromatic ethers and thioethers by means of the Mitsunobu reaction is described, for example, in R. C. Larock, *Comprehensive Organic Transformations* 448 (VCH, 1989), in F.A.

Carey and R. J. Sundberg, *Advanced Organic Chemistry*, Part B 153-4 (Plenum, 2001), and in *Org. React.*, 1992, 42, 335. The phenol and the alcohol or thiol component are reacted together in an aprotic solvent such as, for example, tetrahydrofuran, in the presence of a dialkyl azodicarboxylate and a triarylphosphine, to afford the ether or thioether products. The procedure is also described in *Org. React.*, 1992, 42, 335-656.

Using the above procedures, but employing different hydroxy-substituted hydrazines, and/or different bromo- or mercapto-substituted phosphonates, products analogous to 139.5 and 139.7 are obtained.

Example 140



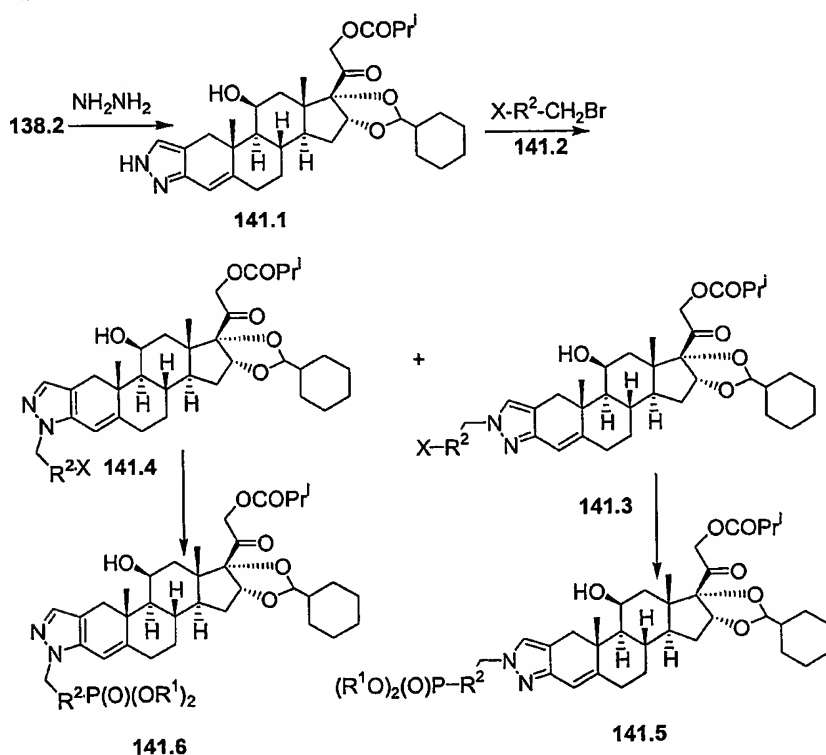
The ketoaldehyde, 138.2, is reacted, as described above, with 4-aminophenyl hydrazine, 140.1, (Epsilon) to produce the pyrazoles, 140.2 and 140.3. The 2'-substituted isomer, 140.2, is coupled, as described above, with a dialkyl phosphonoacetic acid, 140.4, (Aldrich) and dicyclohexyl carbodiimide, to give the amide phosphonate, 140.5.

Alternatively, the 1'-substituted pyrazole, 140.3, is reacted with a dialkyl 3-hydroxypropyl phosphonate, 140.6, (*Zh. Obschei. Khim.*, 1973, 43, 2364), and carbonyl diimidazole to prepare the carbamate phosphonate, 140.7. The preparation of carbamates is described in *Comprehensive Organic Functional*

Group Transformations, Vol. 6, 416ff (A. R. Katritzky, ed., Pergamon, 1995) and in S. R. Sandler and W. Karo, *Organic Functional Group Preparations* 260ff (Academic Press, 1986). In the procedure, the amine is reacted in an inert aprotic solvent such as dichloromethane or tetrahydrofuran, with phosgene or a functional equivalent thereof, such as carbonyl diimidazole, triphosgene, pentafluorophenyl carbonate and the like, to afford the corresponding activated acylamine. The latter compound is then reacted with an alcohol to yield the carbamate.

Using the above procedures, but employing, in place of the 4-amino-phenyl hydrazine, **140.1**, different amino-substituted hydrazines, and/or different dialkyl carboxy or hydroxy-substituted phosphonates, products analogous to the compounds, **140.5** and **140.7** are obtained.

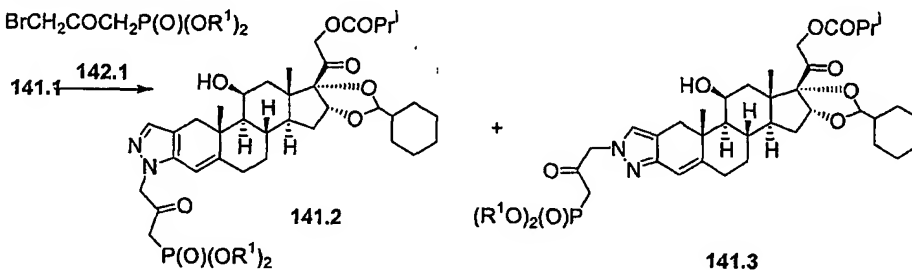
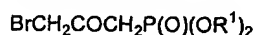
Example 141



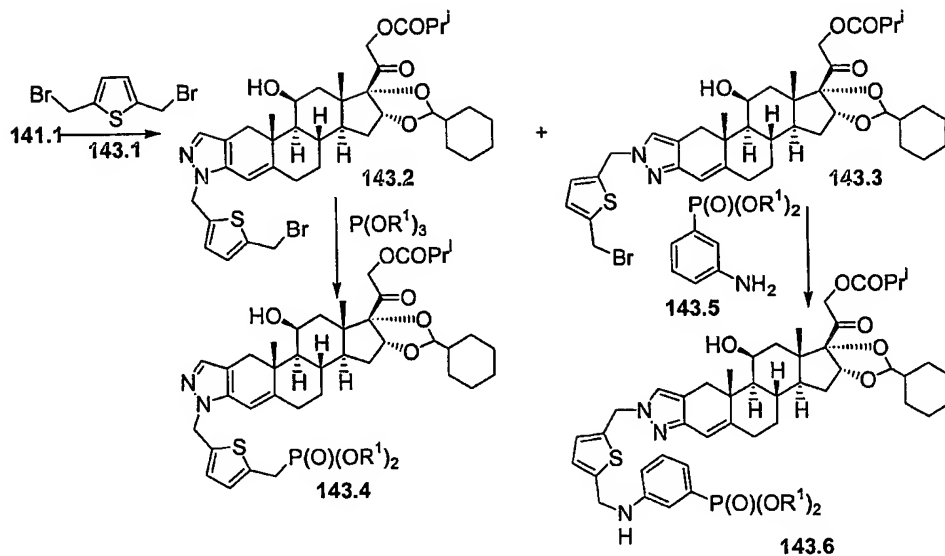
The ketoaldehyde, **138.2**, is reacted with hydrazine, to afford the pyrazole derivative, **141.1**. The reaction of steroidal 2-formyl-3-ketones with hydrazine is described in *J. Am. Chem. Soc.*, 1964, 86, 1520. The reaction is performed in acetic acid at ambient temperature. The pyrazole product is then

reacted with a bromomethyl compound, 141.2, in which R^2 and X are as defined above, or a reactive bromoheteroaromatic reagent, to yield the alkylation products, 141.3 and 141.4. The alkylation of substituted pyrazoles is described, for example, in T. L. Gilchrist, *Heterocyclic Chemistry* 309 (Longman, 1992). The reaction is performed between equimolar amounts of the substrates in a polar solvent such as dimethylformamide or tetrahydrofuran, in the presence of a base such as dimethylaminopyridine, lithium hexamethyldisilazide and the like. The products, 141.3 and 141.4, are, except in cases where X is dialkylphosphono, converted into the phosphonates, 141.5 and 141.6, using the procedures described herein.

Example 142



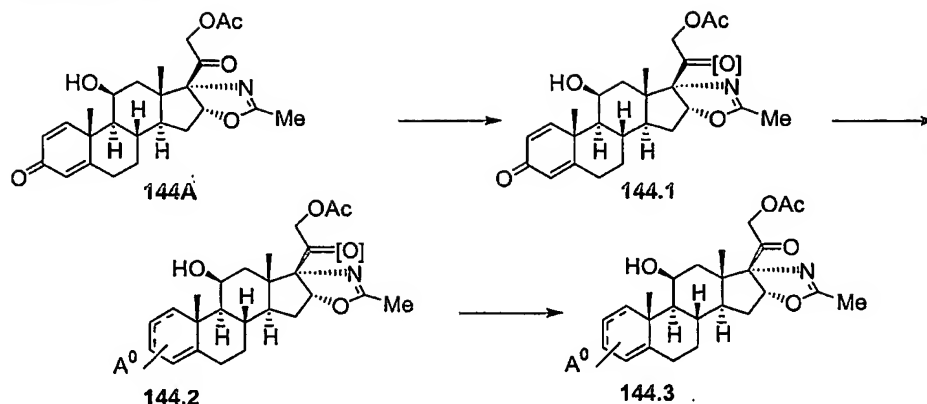
The pyrazole, 141.1, is reacted, as described above, with a dialkyl acetonyl phosphonate, 142.1, (*Tetrahedron Lett.*, 1978, 34, 649) to give the pyrazoles, 141.2 and 141.3.

Example 143

The pyrazole, **141.1**, is reacted in tetrahydrofuran solution, with 2,5-bis(bromomethyl)thiophene, **143.1**, (*Tetrahedron Lett.*, **1999**, 55, 4709) and potassium hexamethyl disilazide, to give the alkylation products, **143.2** and **143.3**. The 2'-substituted isomer, **143.2**, is then reacted, in a Arbuzov reaction, with a trialkyl phosphite to yield the phosphonate, **143.4**. The Arbuzov reaction is described in *Handb. Organophosphorus Chem.*, 1992, 115. In this procedure, in which a bromo substituent is converted into the corresponding phosphonate, the substrate is heated at from about 60°C to about 160°C with a five to fifty-fold molar excess of a trialkyl phosphite, to effect the transformation.

The 2''-substituted pyrazole, **143.3**, is reacted at 70°C in dimethylformamide solution with one molar equivalent of a dialkyl 3-aminophenyl phosphonate, **143.5**, and cesium carbonate, to give the amine phosphonate, **143.6**.

Using the above procedures, but employing different dibromides, and/or different amino-substituted phosphonates, products analogous to **143.4** and **143.6** are obtained.

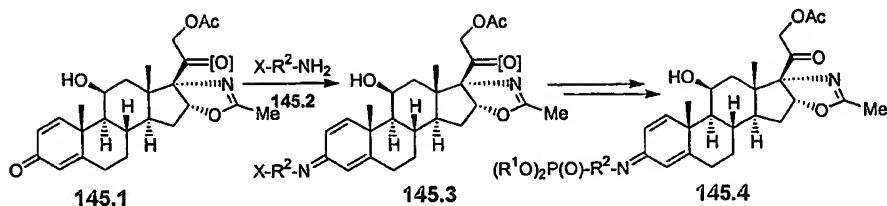
Example 144

A protection-deprotection sequence in which the 20-ketone group of Deflazacort, 144A, (U.S. Patent No. 3,436,389) is protected to afford the derivative, 144.1, as shown. The ketone is protected, for example, by conversion to the cyclic ethylene ketal, by reaction in toluene solution at reflux temperature with ethylene glycol and an acid catalyst, as described in *J. Am. Chem. Soc.*, 1955, 77, 1904. Deprotection is effected by reaction with pyridinium tosylate in aqueous acetone, as described in *J. Chem. Soc. Chem. Comm.*, 1987, 1351.

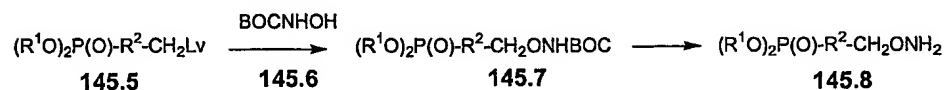
Alternatively, the 20-ketone is protected by conversion to the N, N-dimethylhydrazone. The dimethyl hydrazone is prepared by the reaction of the ketone, 144A, with N,N-dimethylhydrazine in ethanol-acetic acid, as described in *Org. Syn.*, 1970, 50, 102. The group is removed by treatment with sodium acetate and acetic acid in aqueous tetrahydrofuran, as described in *J. Am. Chem. Soc.*, 1979, 101, 5841.

Alternatively, the 20-ketone is protected as the diethylamine adduct. In this procedure, the substrate, 144A, is reacted with titanium tetrakis(diethylamide), as described in *J. Chem. Soc. Chem. Comm.*, 1983, 406, to afford the adduct. The ketone is deprotected by reaction with water in an aqueous organic solvent.

The protected compound, 144.1, is then converted into the phosphonate-containing analog, 144.2, using the procedures described below, and the protecting group is then removed, as described above, to give the phosphonate, 144.3.

Example 145

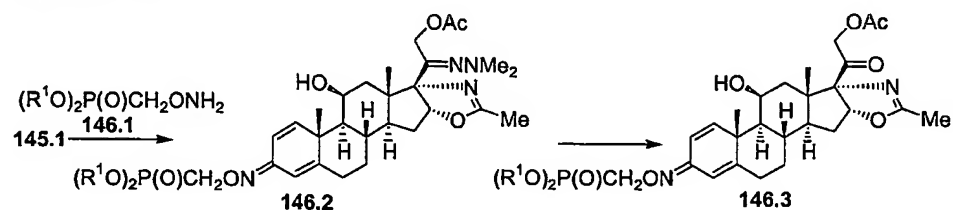
The protected derivative, **145.1**, is reacted with an amine or hydroxylamine, **145.2**, in which R² is an alkyl, alkenyl, cycloalkyl or cycloalkenyl group, optionally incorporating a heteroatom O, S or N, or a functional group such as an amide, ester, oxime, sulfoxide or sulfone etc, or an optionally substituted aryl, heteroaryl or aralkyl group, optionally incorporating a heteroatom O, S or N, and X is either a phosphonate group or a group which is subsequently converted into a phosphonate-containing substituent. For example, X is dialkylphosphono, bromo, hydroxy, amino, carboxy and the like. The reaction is conducted between equimolar amounts of the reactants in an aprotic solvent such as pyridine or xylene, or in an alcoholic solvent such as ethanol, optionally in the presence of an acid catalyst, to give the imine or oxime, **145.3**. The preparation of oximes of steroidal 3-ketones is described in *Anal. Bioch.*, **1978**, *86*, 133 and in *J. Mass. Spectrom.*, **1995**, *30*, 497. The protecting group is then removed, as described herein, to afford the 20-keto phosphonate product, **145.4**.

Example 145A

The preparation of hydroxylamine ethers incorporating a phosphonate group is illustrated herein. In this procedure, a phosphonate, **145.5**, in which Lv is a leaving group such as bromo or trifluoromethylsulfonyloxy, is reacted with BOC-hydroxylamine, **145.6**, (Aldrich) to produce the ether, **145.7**. The reaction is conducted between equimolar amounts of the reactants in a polar solvent such as dimethylformamide or tetrahydrofuran, in the presence of a base such as potassium hydroxide or dimethylaminopyridine. Deprotection, for example by treatment with trifluoroacetic acid, then gives the hydroxylamine ether, **145.8**.

The above procedure is also employed for the preparation of substituted hydroxylamines which are precursors to phosphonates.

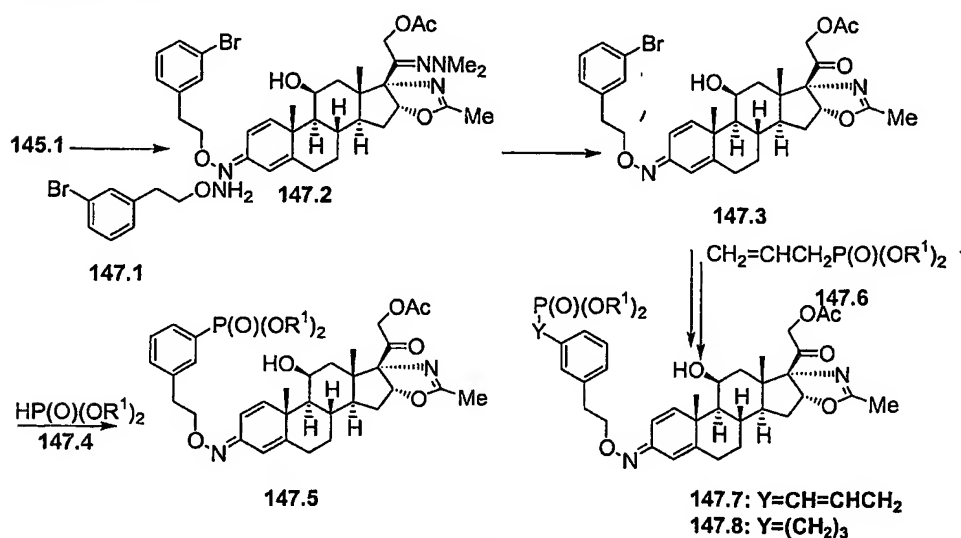
Example 146



The substrate, 145.1, in which the 20-ketone is protected as the dimethyl hydrazone derivative, is reacted with a dialkyl phosphonomethyl hydroxylamine, 146.1, prepared as described above from a dialkyl trifluoromethylsulfonyloxymethyl phosphonate (*Tetrahedron Lett.*, 1986, 27, 1477) and BOC-hydroxylamine, to afford the oxime, 146.2. Deprotection, as described in Example 144, then affords the 20-keto phosphonate, 146.3. The oxime forming reaction is performed at ambient temperature in ethanol-acetic acid solution between equimolar amounts of the reactants.

Using the above procedures, but employing, in place of the hydroxylamine ether, 146.1, different oxime ethers, 145.2, the corresponding products, 145.4 are obtained.

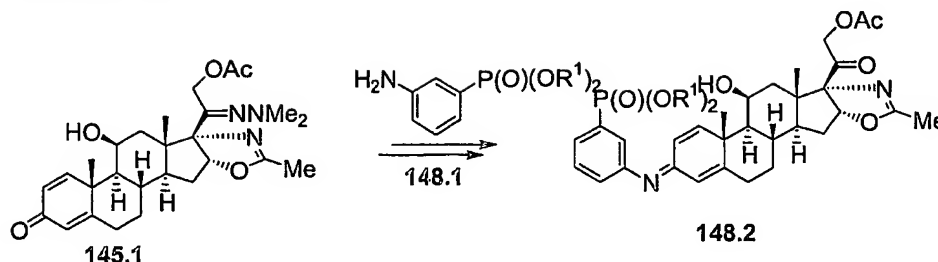
Example 147



The dienone, 145.1, in which the 20-ketone is protected as the dimethyl hydrazone, is reacted, as described above, with O-(3-bromophenylethoxy)-hydroxylamine, 147.1, prepared as described above from 3-bromophenylethyl bromide (French Patent FR 1481052), and BOC-protected hydroxylamine, 145.6, to give the oxime, 147.2. The protecting group is then removed to yield the 20-keto product, 147.3. The latter product is then reacted, in the presence of a palladium catalyst, with a dialkyl phosphate, 147.4, to afford the phosphonate, 147.5. The preparation of arylphosphonates by means of a coupling reaction between aryl bromides and dialkyl phosphites is described in *J. Med. Chem.*, 1992, 35, 1371. The reaction is performed at ca. 100°C in an inert solvent such as toluene, in the presence of a base such as triethylamine and a catalytic amount of tetrakis(triphenylphosphine)palladium(0).

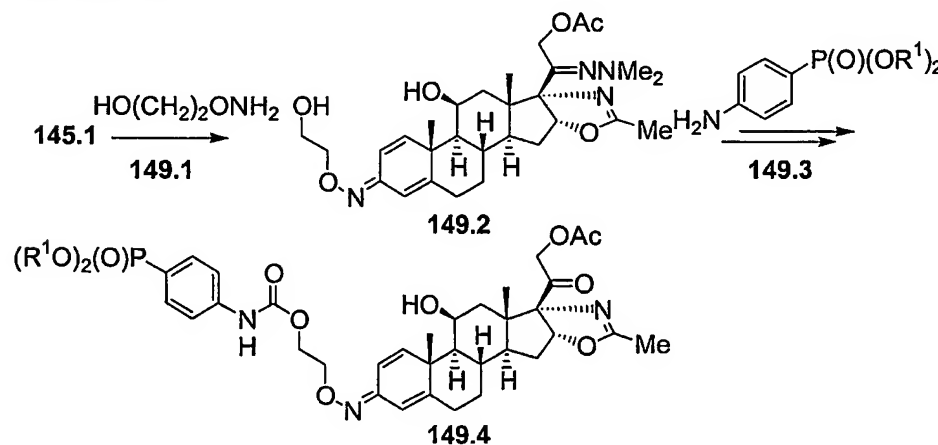
Alternatively, the bromo compound, 147.3, is coupled with a dialkyl propenyl phosphonate, 147.6, (Aldrich) to afford the phosphonate, 147.7. The coupling of aryl halides with olefins by means of the Heck reaction is described, for example, in F. A. Carey and R. J. Sundberg, *Advanced Organic Chemistry* 503ff (Plenum, 2001) and in *Acc. Chem. Res.*, 1979, 12, 146. The aryl bromide and the olefin are coupled in a polar solvent such as dimethylformamide or dioxan, in the presence of a palladium(0) catalyst such as tetrakis(triphenylphosphine)palladium(0) or palladium(II) catalyst such as palladium(II) acetate, and optionally in the presence of a base such as triethylamine or potassium carbonate. Optionally, the styrenoid double bond present in the product, 147.7, is reduced, for example by reaction with diimide, to produce the saturated analog, 147.8. The reduction of olefinic bonds is described in R. C. Larock, *Comprehensive Organic Transformations* 6ff (VCH 1989). The transformation is effected by means of catalytic hydrogenation, for example using a palladium on carbon catalyst and hydrogen or a hydrogen donor, or by the use of diimide or diborane.

Using the above procedures, but employing, in place of the bromophenylethyl reagent, 147.1, different bromo-substituted aryl or heteroaryl alkoxy hydroxylamines, and/or different dialkyl alkenyl phosphonates, products analogous to the compounds, 147.5, 147.7 and 147.8 are obtained.

Example 148

The substrate, **145.1**, in which the 20-ketone is protected as the dimethylhydrazone, is reacted with a dialkyl 3-aminophenyl phosphonate, **148.1**, (*J. Med. Chem.*, **1984**, *27*, 654), to give, after deprotection, the imine product, **148.2**. The imine forming reaction is conducted in a hydrocarbon solvent such as toluene or xylene, at reflux temperature, in the presence of a basic catalyst such as sodium methoxide, or an acid catalyst such as p-toluenesulfonic acid, under azeotropic conditions.

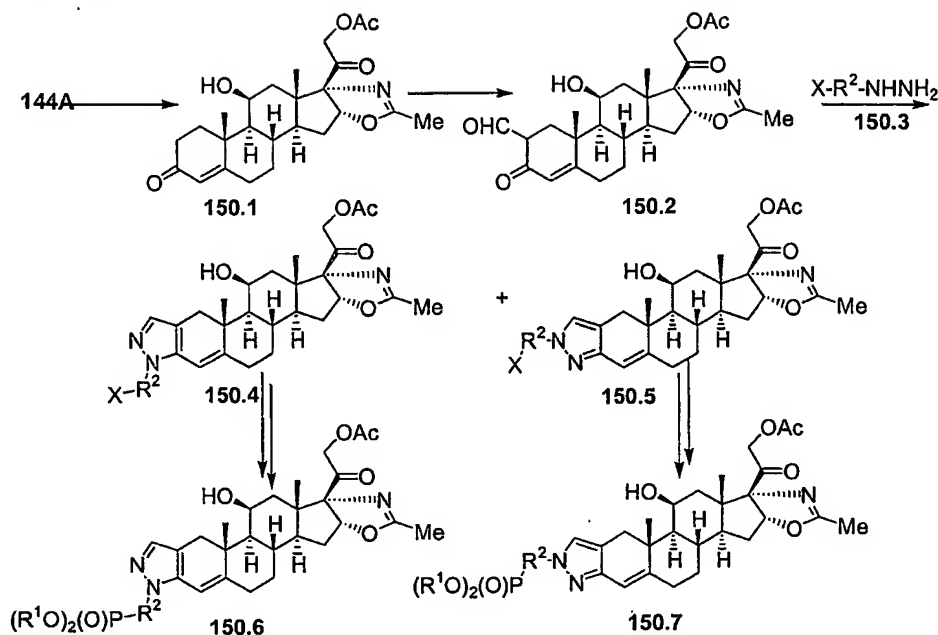
Using the above procedures, but employing, in place of the 3-aminophenyl phosphonate, **148.1**, different amino-substituted aryl or heteroaryl phosphonates, products analogous to **148.2** are obtained.

Example 149

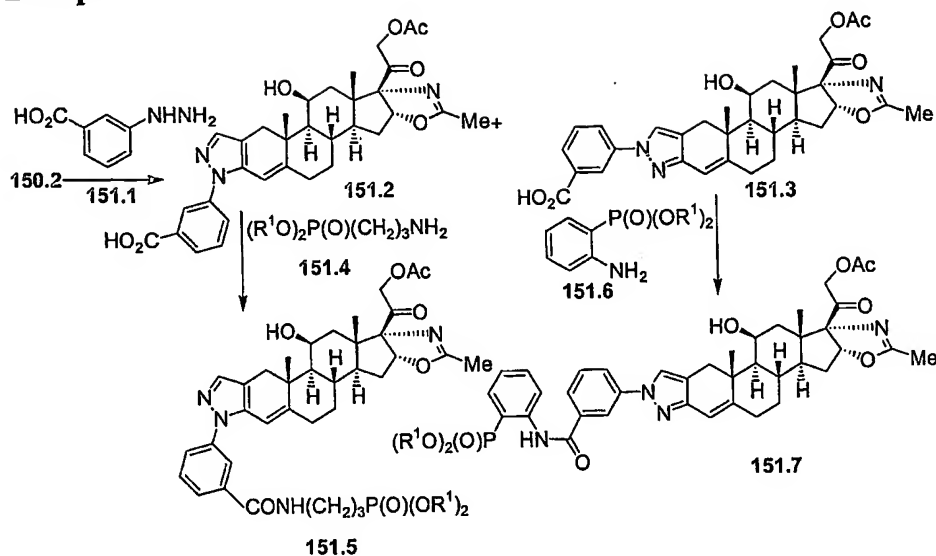
The dienone, **145.1**, in which the 20-ketone is protected as the dimethylhydrazone, is reacted with O-(2-hydroxyethyl)hydroxylamine, **149.1**, (*J. Chem. Soc. Chem. Comm.*, **1986**, 903) to yield the oxime, **149.2**. The reaction of

steroidal 1,4-dien-3-ones with substituted hydroxylamines is described in *J. Steroid Bioch.*, **1976**, *7*, 795; the reaction is performed between equimolar amounts of the reactants in a polar organic solvent such as pyridine or methanol, optionally in the presence of acetic acid or sodium acetate. The product, **149.2**, is then coupled with a dialkyl 4-aminophenyl phosphonate, **149.3**, (Epsilon) and carbonyl diimidazole, to yield, after deprotection, the carbamate oxime, **149.4**. The preparation of carbamates is described in *Comprehensive Organic Functional Group Transformations*, Vol. 6, 416ff (A. R. Katritzky, ed., Pergamon, 1995) and in S. R. Sandler and W. Karo, *Organic Functional Group Preparations* 260ff (Academic Press, 1986). In the procedure, the amine is reacted in an inert aprotic solvent such as dichloromethane or tetrahydrofuran, with phosgene or a functional equivalent thereof, such as carbonyl diimidazole, triphosgene, pentafluorophenyl carbonate and the like, to afford the corresponding activated acylamine. The latter compound is then reacted with an alcohol to yield the carbamate.

Using the above procedures, but employing, in place of the hydroxy-substituted hydroxylamine, **149.1**, different hydroxy-substituted hydroxylamines, and/or different amino-substituted phosphonates, the products analogous to **149.4** are obtained.

Example 150

The dienone, **144A**, is reduced to afford the 1,2-dihydro product, **150.1**. The catalytic hydrogenation reaction is effected by the use of tris(triphenylphosphine)rhodium (I) chloride, for example, as described in *J. Med. Chem.*, **2001**, *44*, 602. The product is then reacted with ethyl formate and a base such as sodium hydride, in an inert solvent such as toluene or dimethylformamide, as described in *J. Am. Chem. Soc.*, **1964**, *86*, 1520, to afford the 2-formyl product, **150.2**. This compound is then reacted with an alkyl, aralkyl, aryl or heteroaryl hydrazine, **150.3**, in which the substituent X is either a phosphonate group or a group which is subsequently transformed into a phosphonate-containing substituent. For example, X is dialkylphosphono, bromo, hydroxy, amino, carboxyl and the like. The reaction yields the isomeric 2'- and 1'-aryl pyrazoles, **150.4** and **150.5**. The pyrazole-forming reaction is performed between equimolar amounts of the reactants in an acidic solvent such as acetic acid, as described in *J. Am. Chem. Soc.*, **1964**, *86*, 1520. The pyrazoles, **150.4** and **150.5**, are then transformed, for example by the procedures described herein, into the phosphonates, **150.6** and **150.7**.

Example 151

The ketoaldehyde, **150.2**, is reacted, as described above, with 3-carboxyphenyl hydrazine, **151.1**, (Apin) to give the pyrazoles, **151.2** and **151.3**. The 2'-substituted isomer, **151.2**, is then coupled in dimethylformamide solution at ambient temperature with a dialkyl 3-aminopropyl phosphonate, **151.4**, (Acros) and dicyclohexyl carbodiimide, to yield the amide phosphonate, **151.5**. The preparation of amides from carboxylic acids and derivatives is described, for example, in S.R. Sandler and W. Karo, *Organic Functional Group Preparations* 274 (Academic Press, 1986), and R. C. Larock, *Comprehensive Organic Transformations* 972ff (VCH, 1989). The carboxylic acid is reacted with the amine in the presence of an activating agent, such as, for example, dicyclohexylcarbodiimide or diisopropylcarbodiimide, optionally in the presence of, for example, hydroxybenztriazole, N-hydroxysuccinimide or N-hydroxypyridone, in a non-protic solvent such as, for example, pyridine, DMF or dichloromethane, to afford the amide.

Alternatively, the carboxylic acid may first be converted into an activated derivative such as the acid chloride, anhydride, mixed anhydride, imidazolid and the like, and then reacted with the amine, in the presence of an organic base such as, for example, pyridine, to afford the amide.

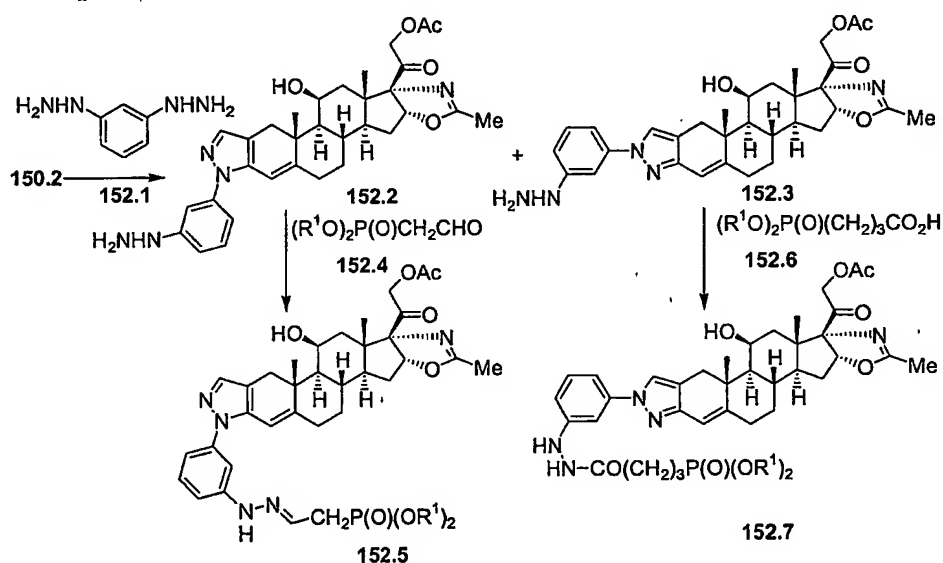
The conversion of a carboxylic acid into the corresponding acid chloride can be effected by treatment of the carboxylic acid with a reagent such as, for

example, thionyl chloride or oxalyl chloride in an inert organic solvent such as dichloromethane, optionally in the presence of a catalytic amount of dimethylformamide.

The isomeric pyrazole, **151.3**, is reacted, as described above, with a dialkyl 2-aminophenyl phosphonate, **151.6**, (Acros) to yield the amide phosphonate, **151.7**.

Using the above procedures, but employing different carboxy-substituted hydrazines, and/or different amino-substituted phosphonates, the products analogous to **151.5** and **151.7** are obtained.

Example 152

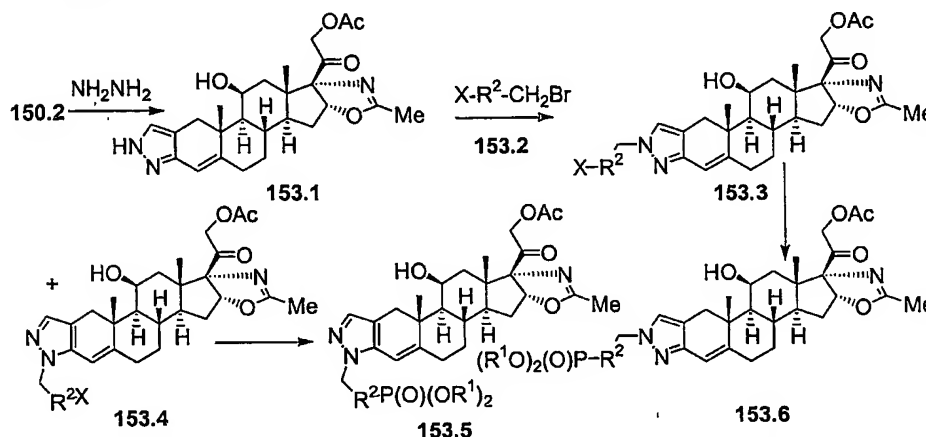


The ketoaldehyde, **150.2**, is reacted, as described above, with 1,3-bis(hydrazino)benzene, **152.1**, (*Bull. Soc. Chim. Fr.*, 1975, 1371) to produce the pyrazoles, **152.2** and **152.3**. The 2'-substituted isomer, **152.2**, is reacted in tetrahydrofuran solution at ambient temperature with one molar equivalent of a dialkylphosphono acetaldehyde (Aurora), to give the hydrazone phosphonate, **152.5**.

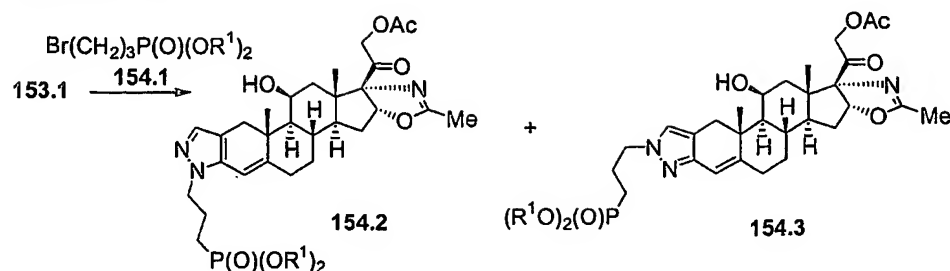
Alternatively, the 1'-substituted pyrazole, **152.3**, is coupled, as described above, with a dialkylphosphono butyric acid, **152.6**, (Epsilon) and dicyclohexyl carbodiimide to prepare the phosphonate, **152.7**.

Using the above procedures, but employing, in place of the 1,3-bis(hydrazino)phenyl hydrazine, **152.1**, different bis hydrazines, and/or different dialkyl formyl or carboxy-substituted phosphonates, the products analogous to the compounds, **152.5** and **152.7** are obtained.

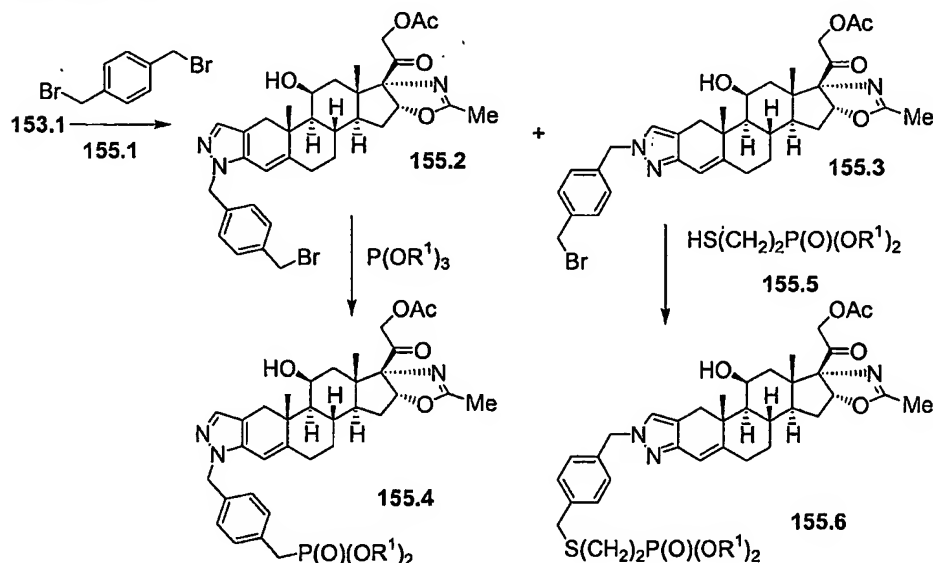
Example 153



The ketoaldehyde, **150.2**, is reacted with hydrazine to afford the pyrazole derivative, **153.1**. The reaction of steroidal 2-formyl-3-ketones with hydrazine is described in *J. Am. Chem. Soc.*, **1964**, *86*, 1520. The reaction is performed in acetic acid at ambient temperature. The pyrazole product is then reacted with a bromomethyl compound **153.2**, in which R² and X are as defined above, or a reactive bromoheteroaromatic reagent, to yield the alkylation products, **153.3** and **153.4**. The alkylation of substituted pyrazoles is described, for example, in T. L. Gilchrist, *Heterocyclic Chemistry* 309 (Longman, 1992). The reaction is performed between equimolar amounts of the substrates in a polar solvent such as dimethylformamide or tetrahydrofuran, in the presence of a base such as dimethylaminopyridine, lithium hexamethyldisilazide and the like. The products, **153.3** and **153.4**, are, except in cases where X is dialkylphosphono, converted into the phosphonates, **153.5** and **153.6**, using the procedures described herein.

Example 154

The pyrazole, **153.1**, is reacted in dimethylformamide solution at 70°C with one molar equivalent of a dialkyl bromopropyl phosphonate, **154.1**, (Synthelec) and cesium carbonate, to give the pyrazoles, **154.2** and **154.3**.

Example 155

The pyrazole, **153.1**, is reacted in tetrahydrofuran solution with 1,4-bis(bromomethyl)benzene, **155.1**, and potassium hexamethyl disilazide, to give the alkylation products, **155.2** and **155.3**. The 2'-substituted isomer, **155.2**, is then reacted, in an Arbuzov reaction, with a trialkyl phosphite to yield the phosphonate, **155.4**. The Arbuzov reaction is described in *Handb.*

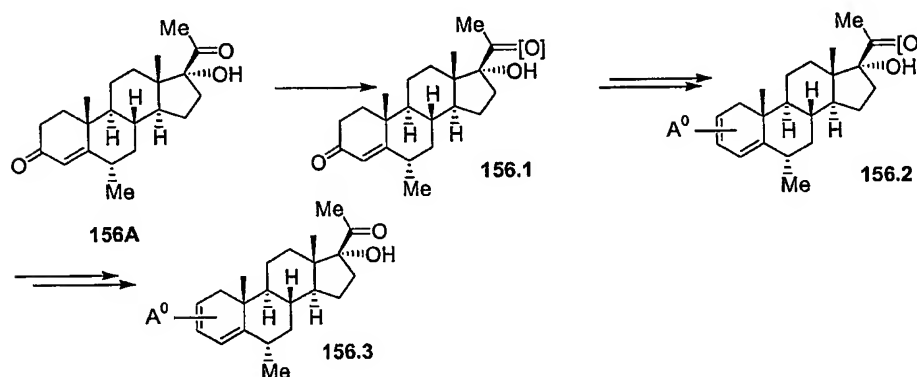
Organophosphorus Chem., 1992, 115. In this procedure, in which a bromo substituent is converted into the corresponding phosphonate, the substrate is

heated at from about 60°C to about 160°C with a five to fifty-fold molar excess of a trialkyl phosphite, to effect the transformation.

The 2'-substituted pyrazole, **155.3**, is reacted at 70°C in dimethyl-formamide solution with one molar equivalent of a dialkyl mercaptoethyl phosphonate, **155.5**, (*Zh. Obschei. Khim.*, **1973**, *43*, 2364) and cesium carbonate, to give the thioether phosphonate, **155.6**.

Using the above procedures, but employing different dibromides, and/or different mercapto-substituted phosphonates, products analogous to **155.4** and **155.6** are obtained.

Example 156



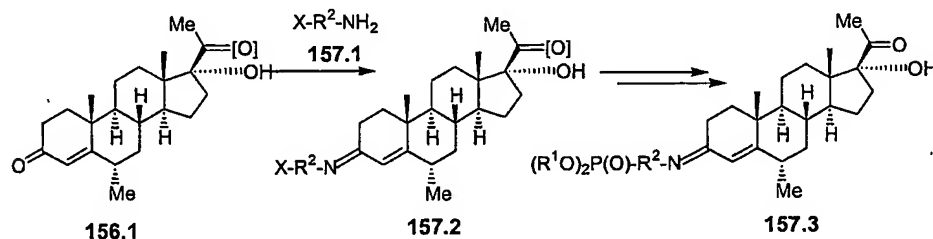
Medroxyprogesterone, **156A**, (U.S. Patent Nos. 3,043,832, 3,061,616, and 3,377,364) is protected to afford the derivative, **156.1**. The ketone is protected, for example, by conversion to the cyclic ethylene ketal, by reaction in toluene solution at reflux temperature with ethylene glycol and an acid catalyst, as described in *J. Am. Chem. Soc.*, **1955**, *77*, 1904. Deprotection is effected by reaction with pyridinium tosylate in aqueous acetone, as described in as described in *J. Chem. Soc. Chem. Comm.*, **1987**, 1351.

Alternatively, the 20-ketone is protected by conversion to the N, N-dimethylhydrazone. The dimethyl hydrazone is prepared by the reaction of the ketone, **156A**, with N,N-dimethylhydrazine in ethanol-acetic acid, as described in *Org. Syn.*, **1970**, *50*, 102. The group is removed by treatment with sodium acetate and acetic acid in aqueous tetrahydrofuran, as described in *J. Am. Chem. Soc.*, **1979**, *101*, 5841.

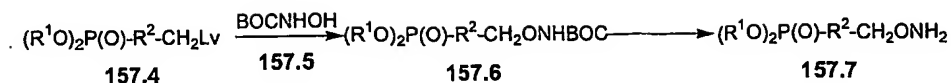
Alternatively, the 20-ketone is protected as the diethylamine adduct. In this procedure, the substrate, **156A**, is reacted with titanium tetrakis(diethylamide), as described in *J. Chem. Soc. Chem. Comm.*, **1983**, 406, to afford the adduct. The ketone is deprotected by reaction with water in an aqueous organic solvent.

The protected compound, **156.1**, is then converted into the phosphonate-containing analog, **156.2**, using the procedures described below, and the protecting group or groups are then removed, as described above, to give the phosphonate, **156.3**.

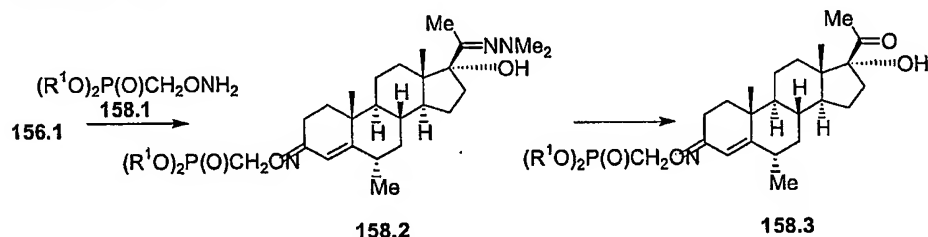
Example 157



The ketone-protected derivative, **156.1**, is reacted with a hydroxylamine or amine, **157.1**, in which R² is an alkyl, alkenyl, cycloalkyl or cycloalkenyl group, optionally incorporating a heteroatom O, S or N, or a functional group such as an amide, ester, oxime, sulfoxide or sulfone, etc., or an optionally substituted aryl, heteroaryl or aralkyl group, optionally incorporating a heteroatom O, S or N, and X is either a phosphonate group or a group which is subsequently converted into a phosphonate-containing substituent. For example, X is dialkylphosphono, bromo, hydroxy, amino, carboxy and the like. The reaction is conducted between equimolar amounts of the reactants in an aprotic solvent such as pyridine or xylene, or in an alcoholic solvent such as ethanol, optionally in the presence of an acid catalyst, to give the oxime, **157.2**. The preparation of oximes of steroidal 3-ketones is described in *Anal. Bioch.* **1978**, 86, 133 and in *J. Mass. Spectrom.* **1995**, 30, 497. The protecting group is then removed, as described in Example 156, to afford the 20-keto phosphonate product, **157.3**.

Example 157A

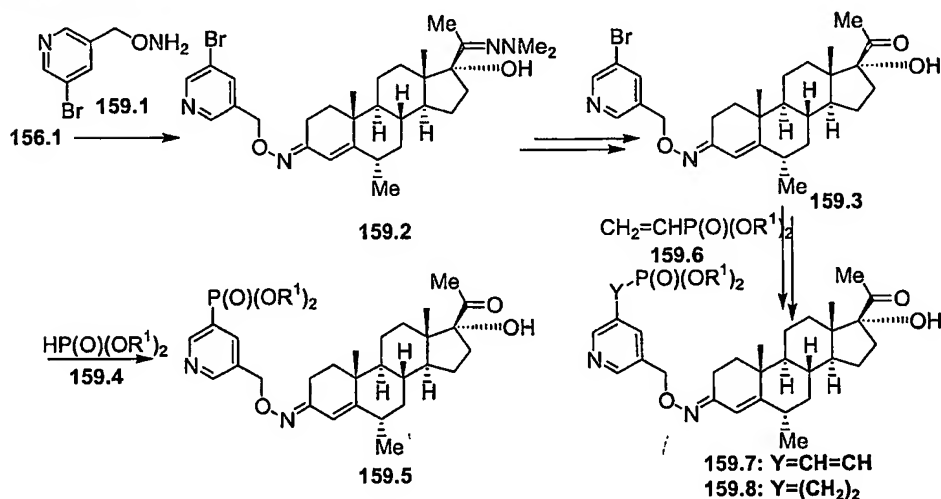
The preparation of hydroxylamine ethers incorporating a phosphonate group is illustrated herein. A phosphonate, **157.4**, in which Lv is a leaving group such as bromo or trifluoromethylsulfonyloxy, is reacted with BOC-hydroxylamine, **157.5**, (Aldrich) to produce the ether, **157.6**. The reaction is conducted between equimolar amounts of the reactants in a polar solvent such as dimethylformamide or tetrahydrofuran, in the presence of a base such as potassium hydroxide or dimethylaminopyridine. Deprotection, for example by treatment with trifluoroacetic acid, then gives the hydroxylamine ether, **157.7**. The above procedure is also employed for the preparation of substituted hydroxylamines which are precursors to phosphonates.

Example 158

The preparation of phosphonates, in which the phosphonate is attached by means of an iminoxy group is illustrated herein. In this procedure, the substrate, **156.1**, in which the 20-ketone is protected as the dimethyl hydrazone derivative, is reacted with a dialkyl phosphonomethyl hydroxylamine, **158.1**, prepared as described above from a dialkyl trifluoromethylsulfonyloxymethyl phosphonate (*Tetrahedron Lett.*, **1986**, 27, 1477) and BOC-hydroxylamine, to afford the oxime, **158.2**. Deprotection, as described in Example 156, then affords the 20-keto phosphonate, **158.3**. The oxime forming reaction is performed at ambient temperature in ethanol-acetic acid solution between equimolar amounts of the reactants.

Using the above procedures, but employing, in place of the hydroxylamine ether, **158.1**, different oxime ethers, **157.1**, the corresponding products, **157.3** are obtained.

Example 159



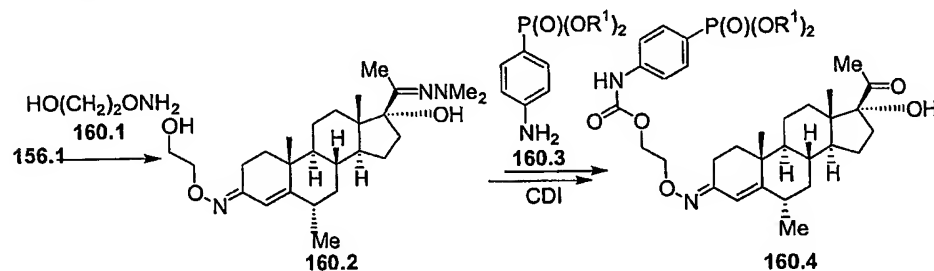
The dienone, **156.1**, in which the 20-ketone is protected as the dimethyl hydrazone, is reacted, as described above, with O-(5-bromo-3-pyridylmethoxy)-hydroxylamine, **159.1**, prepared as described above from 5-bromo-3-bromo-methylpyridine (WO 95/28400) and BOC-protected hydroxylamine, **157.5**, to give the oxime, **159.2**. The protecting group is then removed to yield the 20-keto product, **159.3**. The latter product is then reacted, in the presence of a palladium catalyst, with a dialkyl phosphite, **159.4**, to afford the phosphonate, **159.5**. The preparation of arylphosphonates by means of a coupling reaction between aryl bromides and dialkyl phosphites is described in *J. Med. Chem.*, **1992**, 35, 1371. The reaction is performed at ca. 100°C in an inert solvent such as toluene, in the presence of a base such as triethylamine and a catalytic amount of tetrakis(triphenylphosphine)palladium(0).

Alternatively, the bromo compound, **159.3**, is coupled with a dialkyl vinylphosphonate, **159.6**, (Aldrich) to afford the phosphonate, **159.7**. The coupling of aryl halides with olefins by means of the Heck reaction is described, for example, in F. A. Carey and R. J. Sundberg, *Advanced Organic Chemistry* 503ff (Plenum, 2001) and in *Acc. Chem. Res.*, **1979**, 12, 146. The aryl bromide

and the olefin are coupled in a polar solvent such as dimethylformamide or dioxan, in the presence of a palladium(0) catalyst such as tetrakis(triphenylphosphine)palladium(0) or palladium(II) catalyst such as palladium(II) acetate, and optionally in the presence of a base such as triethylamine or potassium carbonate. Optionally, the styrenoid double bond present in the product, 159.7, is reduced, for example by reaction with diimide, to produce the saturated analog, 159.8. The reduction of olefinic bonds is described in R. C. Larock, *Comprehensive Organic Transformations* 6ff (VCH 1989). The transformation is effected by means of catalytic hydrogenation, for example using a palladium on carbon catalyst and hydrogen or a hydrogen donor, or by the use of diimide or diborane.

Using the above procedures, but employing, in place of the bromopyridyl reagent, 159.1, different bromo-substituted aryl or heteroaryl alkoxy hydroxylamines, and/or different dialkyl alkenyl phosphonates, products analogous to the compounds, 159.5, 159.7 and 159.8 are obtained.

Example 160

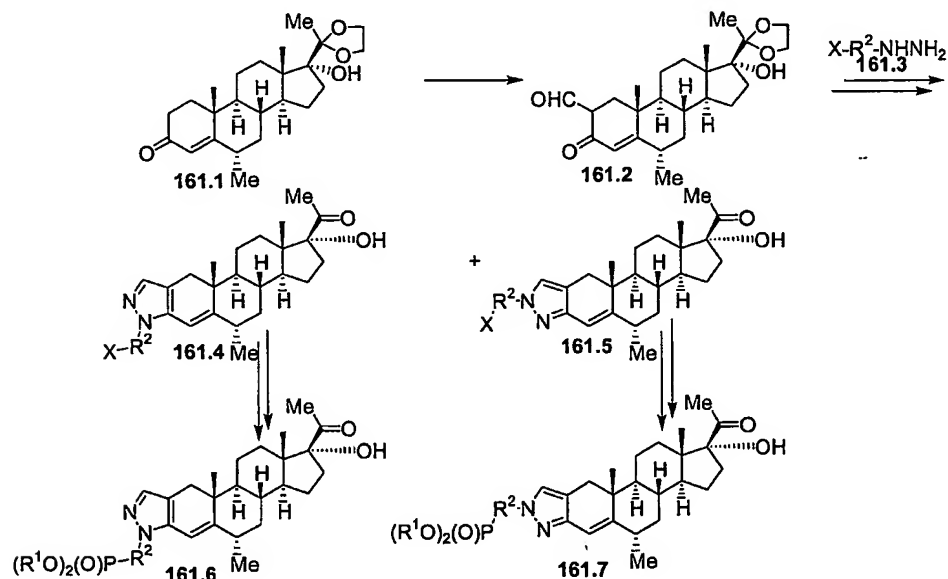


The dienone, 156.1, in which the 20-ketone is protected as the dimethylhydrazone, is reacted with 2-hydroxyethyl hydroxylamine, 160.1, (*J. Chem. Soc. Chem. Comm.*, 1986, 903) to yield the oxime, 160.2. The reaction of unsaturated steroidal ketones with hydroxylamines is described in *J. Steroid Bioch.* 1976, 7, 795; the reaction is performed between equimolar amounts of the reactants in a polar organic solvent such as pyridine or methanol, optionally in the presence of acetic acid or sodium acetate. The product, 160.2, is then coupled with a dialkyl 4-aminophenyl phosphonate, 160.3, (Epsilon) and carbonyl diimidazole, to yield, after deprotection, the carbamate oxime, 160.4. The preparation of carbamates is described in *Comprehensive Organic*

Functional Group Transformations, Vol. 6, 416ff (A. R. Katritzky, ed., Pergamon, 1995) and in S. R. Sandler and W. Karo, *Organic Functional Group Preparations* 260ff (Academic Press, 1986). In the procedure, the amine is reacted in an inert aprotic solvent such as dichloromethane or tetrahydrofuran, with phosgene or a functional equivalent thereof, such as carbonyl diimidazole, triphosgene, pentafluorophenyl carbonate and the like, to afford the corresponding activated acylamine. The latter compound is then reacted with an alcohol to yield the carbamate.

Using the above procedures, but employing, in place of the hydroxy-substituted hydroxylamine, **160.1**, different hydroxy-substituted hydroxylamines, and/or different amino-substituted phosphonates, the products analogous to **160.4** are obtained.

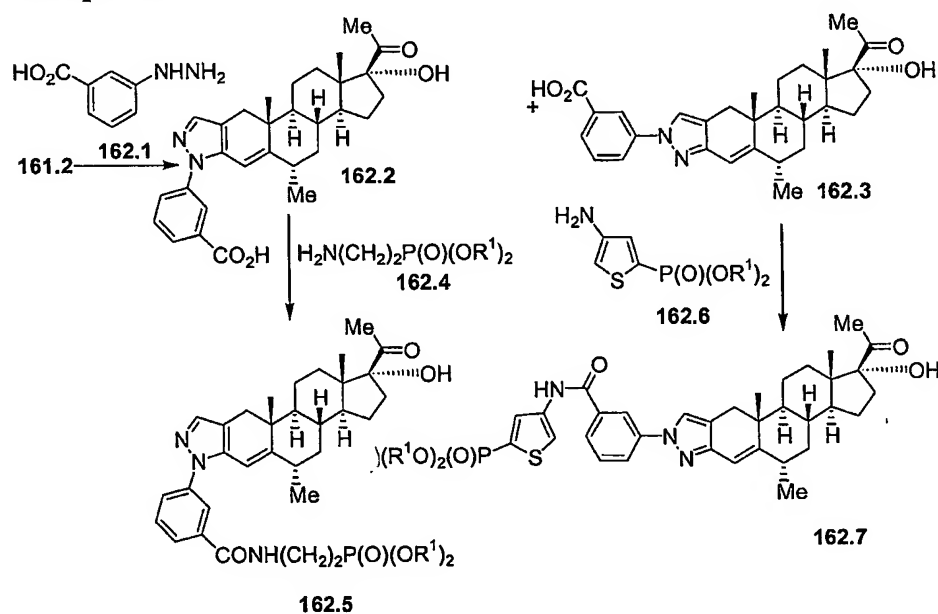
Example 161



The enone, **161.1**, in which the 20-ketone is protected as the cyclic ethylene ketal, is reacted with ethyl formate and a base such as sodium hydride, in an inert solvent such as toluene or dimethylformamide, as described in *J. Am. Chem. Soc.* 86:1520 (1964), to afford the 2-formyl product, **161.2**. This compound is then reacted with an alkyl, aralkyl, aryl or heteroaryl hydrazine, **161.3**, in which the substituent X is either a phosphonate group or a group which is subsequently transformed into a phosphonate-containing substituent. For

example, X is dialkylphosphono, bromo, hydroxy, amino, carboxyl and the like. The reaction yields, after deprotection of the 20-ketone, the isomeric 2'- and 1'-aryl pyrazoles, **161.4** and **161.5**. The pyrazole-forming reaction is performed between equimolar amounts of the reactants in an acidic solvent such as acetic acid, as described in *J. Am. Chem. Soc.*, 1964, 86, 1520. The pyrazoles, **161.4** and **161.5**, are then transformed, for example by the procedures described herein, into the phosphonates, **161.6** and **161.7**.

Example 162



The ketoaldehyde, **161.2**, is reacted, as described above, with 3-carboxyphenyl hydrazine, **162.1**, (Apin) to give the pyrazoles, **162.2** and **162.3**. The 2'-substituted isomer, **162.2**, is then reacted in dimethylformamide solution at ambient temperature with one molar equivalent of a dialkyl 2-aminoethyl phosphonate, **162.4**, (Aldrich) and dicyclohexyl carbodiimide, to give the amide phosphonate, **162.5**. The preparation of amides from carboxylic acids and derivatives is described, for example, in S.R. Sandler and W. Karo, *Organic Functional Group Preparations* 274 (Academic Press, 1968) and R. C. Larock, *Comprehensive Organic Transformations* 972ff (VCH, 1989). The carboxylic acid is reacted with the amine in the presence of an activating agent, such as, for example, dicyclohexylcarbodiimide or diisopropylcarbodiimide, optionally in the presence of, for example, hydroxybenztriazole, N-hydroxysuccinimide or N-

hydroxypyridone, in a non-protic solvent such as, for example, pyridine, DMF or dichloromethane, to afford the amide.

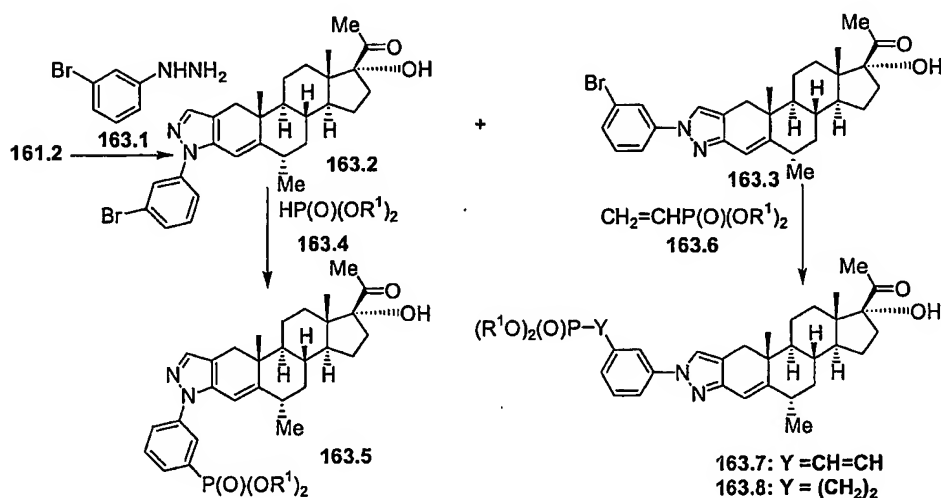
Alternatively, the carboxylic acid may first be converted into an activated derivative such as the acid chloride, anhydride, mixed anhydride, imidazolidine and the like, and then reacted with the amine, in the presence of an organic base such as, for example, pyridine, to afford the amide.

The conversion of a carboxylic acid into the corresponding acid chloride can be effected by treatment of the carboxylic acid with a reagent such as, for example, thionyl chloride or oxalyl chloride in an inert organic solvent such as dichloromethane, optionally in the presence of a catalytic amount of dimethylformamide.

The isomeric pyrazole, 162.3, is reacted, as described above, with one molar equivalent of a dialkyl 4-amino-2-thienyl phosphonate, 162.6, prepared by the palladium catalyzed coupling reaction, as described above, between 4-amino-2-bromothiophene (*Tetrahedron Lett.*, 1987, 43, 3295) and a dialkyl phosphite, to give the amide phosphonate, 162.7.

Using the above procedures, but employing different carboxy-substituted hydrazines, and/or different amino-substituted phosphonates, the products analogous to 162.5 and 162.7 are obtained.

Example 163

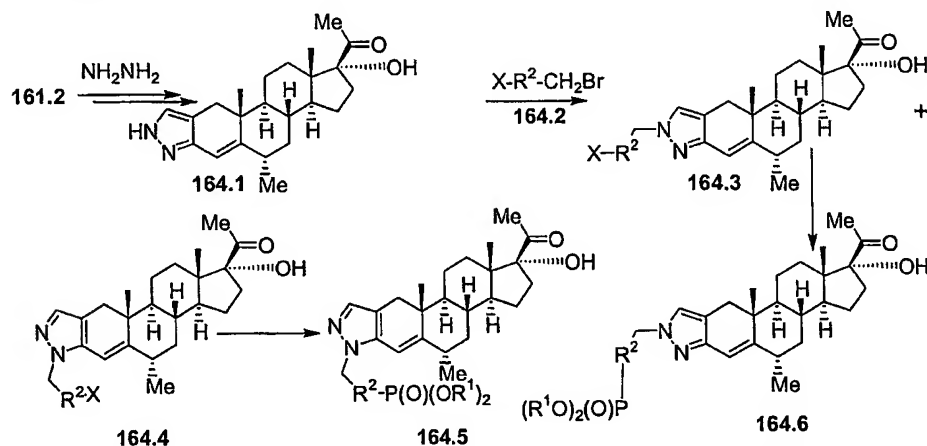


The ketoaldehyde, 161.2, is reacted, as described above, with 3-bromophenyl hydrazine, 163.1, (Fluka) to produce the pyrazoles, 163.2 and 163.3. The 2'-substituted isomer, 163.2, is then coupled, as described above, with a dialkyl phosphite, 163.4, to afford the phosphonate, 163.5.

Alternatively, the 1'-substituted pyrazole, 163.3, is coupled, as described above, with a dialkyl vinylphosphonate, 163.6, (Aldrich) and a palladium catalyst to prepare the vinyl phosphonate, 163.7. Optionally, the product is reduced, as described above, to give the analog, 163.8.

Using the above procedures, but employing, in place of the bromophenyl hydrazine, 163.1, different bromo-substituted hydrazines, and/or different dialkyl alkenyl phosphonates, the products analogous to the compounds, 163.5 and 163.7, 163.8 are obtained.

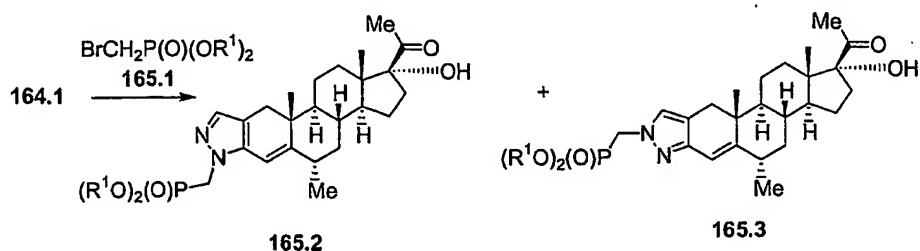
Example 164



The ketoaldehyde, 161.2, is reacted with hydrazine to afford, after deprotection of the 20-ketone, the pyrazole derivative, 164.1. The reaction of steroidal 2-formyl-3-ketones with hydrazine is described in *J. Am. Chem. Soc.*, 1964, 86, 1520. The reaction is performed in acetic acid at ambient temperature. The pyrazole product is then reacted with a bromomethyl compound, 164.2, in which R^2 and X are as defined above, or a reactive bromoheteroaromatic reagent, to yield the alkylation products, 164.3 and 164.4. The alkylation of substituted pyrazoles is described, for example, in T. L. Gilchrist, *Heterocyclic Chemistry*, 309 (Longman, 1992). The reaction is performed between equimolar amounts of the substrates in a polar solvent such as dimethylformamide or

tetrahydrofuran, in the presence of a base such as dimethylaminopyridine, lithium hexamethyldisilazide and the like. The products, **164.3** and **164.4**, are, except in cases where X is dialkylphosphono, converted into the phosphonates, **164.5** and **164.6**, using the procedures described herein.

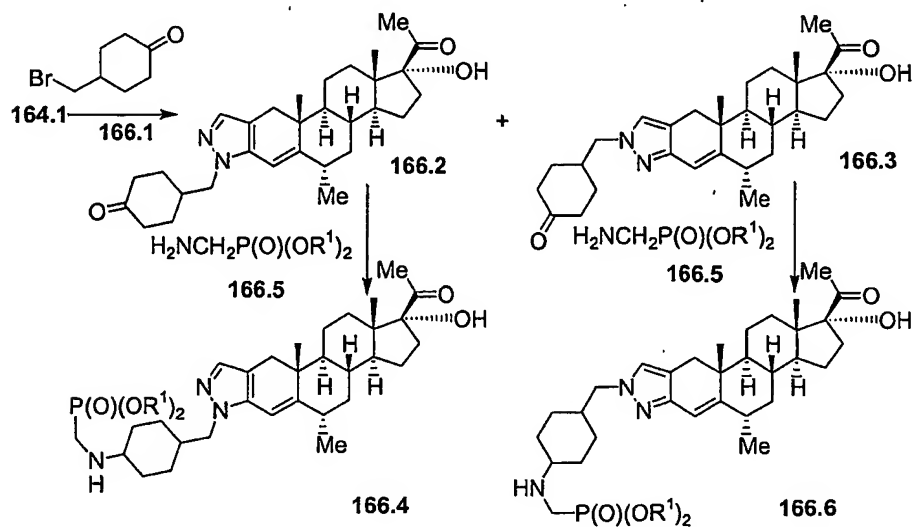
Example 165



The pyrazole, **164.1**, is reacted in dimethylformamide solution at 70°C with one molar equivalent of a dialkyl 4-bromomethyl phosphonate, **165.1**, (Lancaster) and lithium hexamethyl disilazide, to give the pyrazoles, **165.2** and **165.3**.

Using the above procedures, but employing different bromo-substituted phosphonates, the products analogous to **165.2** and **165.3** are obtained.

Example 166

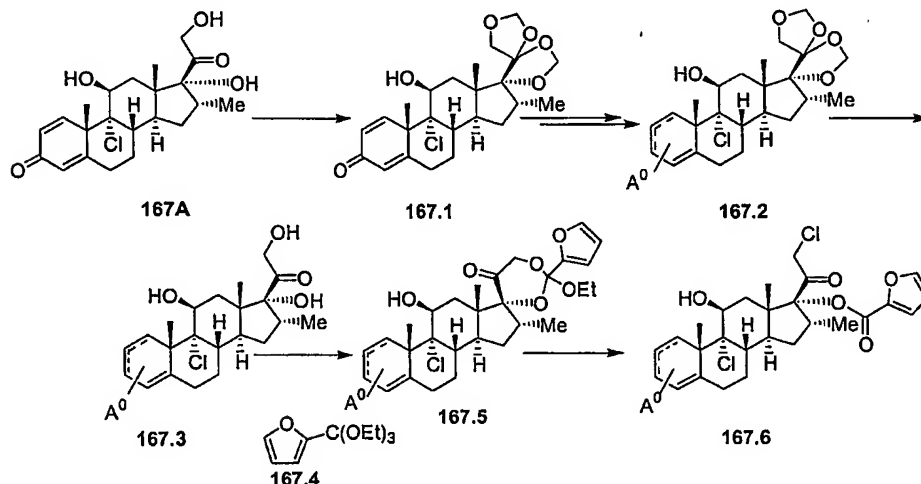


The pyrazole, **164.1**, is reacted in tetrahydrofuran solution with 4-bromomethyl cyclohexanone, **166.1**, (WO 97/37959) and potassium hexamethyl disilazide, to give the alkylation products, **166.2** and **166.3**. The 2'-substituted isomer, **166.2**, is then reacted, in a reductive amination reaction, with a dialkyl aminomethyl phosphonate, **166.5**, (Interchim) and sodium triacetoxy borohydride, to yield the amine phosphonate, **166.4**. The preparation of amines by means of reductive amination procedures is described, for example, in R. C. Larock, *Comprehensive Organic Transformations* 421 (VCH) and in F.A. Carey and R. J. Sundberg, *Advanced Organic Chemistry*, Part B, 269 (Plenum, 2001). In this procedure, the amine component and the aldehyde or ketone component are reacted together in the presence of a reducing agent such as, for example, borane, sodium cyanoborohydride, sodium triacetoxyborohydride or diisobutylaluminum hydride, optionally in the presence of a Lewis acid, such as titanium tetraisopropoxide, as described in *J. Org. Chem.*, 1990, 55, 2552.

The 1'-substituted pyrazole, **166.3**, is converted by the same reaction into the isomeric amine phosphonate, **166.6**.

Using the above procedures, but employing different bromo-substituted aldehydes and ketones, and/or different amino-substituted phosphonates, products analogous to **166.4** and **166.6** are obtained.

Example 167

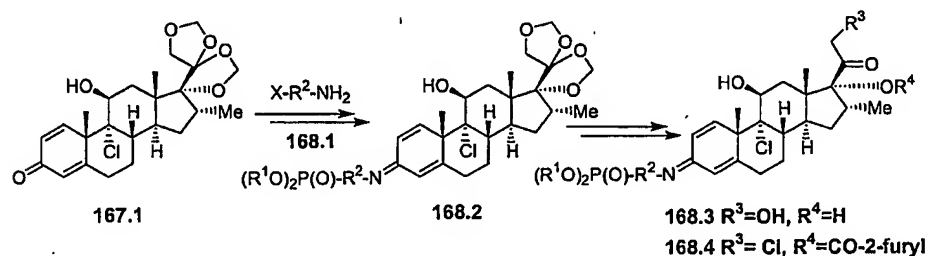


9 α -Chloro-16 α -methyl-11 β ,17 α ,21-trihydroxypregn-1,4-dien-3,21-dione, **167A**, (U.S. Patent No. 4,472,393) is reacted with paraformaldehyde and an acid

catalyst such as hydrochloric acid, as described in *Protective Groups in Organic Synthesis*, by T.W. Greene and P.G.M. Wuts, Wiley, Second Edition 1990, p. 223, to yield the BMD derivative, 167.1. The phosphonate moiety is then introduced, using the procedures described below, to produce the phosphonate ester, 167.2. The BMD moiety is then hydrolyzed, for example by treatment with 50% aqueous acetic acid, as described in *Protective Groups in Organic Synthesis*, by T.W. Greene and P.G.M. Wuts, Wiley, Second Edition 1990, p. 223, to afford the triol, 167.3. The latter compound is then converted into the 17,21-cyclic orthoester, 167.5, using the procedure described in *Chem. Pharm. Bull.*, 1986, 34, 1613. The substrate is reacted in dimethylformamide at 70°C with two molar equivalents of triethyl ortho-2-furoate, 165.4, (*Zh. Org. Khim.*, 1980, 50, 1348) and a catalytic amount of p-toluenesulfonic acid. The product is then reacted with an excess of trimethylsilyl chloride in dimethylformamide at ambient temperature to produce the 21-chloro 17-(2-furoate) product, 167.6.

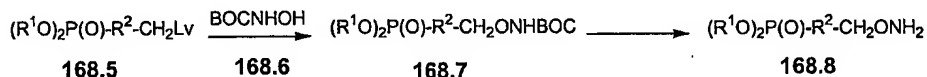
Alternatively, the substrate, 167.3, is converted into the product, 167.6, by means of the method described in *J. Med. Chem.*, 1987, 30, 1581. In this procedure, the 21-hydroxy group is activated by conversion to the 21-mesylate, by reaction with mesyl chloride in pyridine; the mesylate group is then displaced to yield the 21-chloro intermediate, by reaction with lithium chloride in dimethylformamide, and the 17-hydroxyl group is esterified to give the 21-chloro-17-(2-furoate) derivative, 167.6. The selective acylation of the 17 α -hydroxyl group in the presence of an 11 β hydroxyl group is described in *J. Med. Chem.*, 1987, 30, 1581.

Example 168



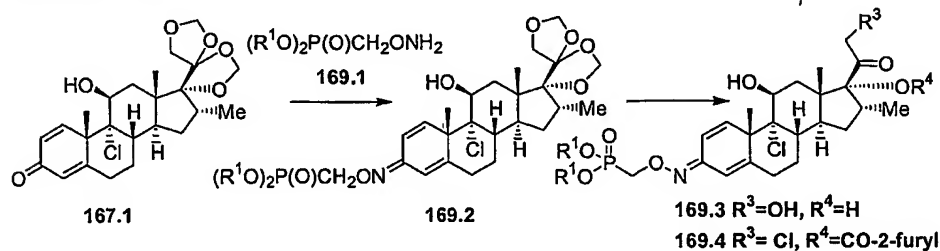
The BMD-protected derivative, 167.1, is reacted with an amine or hydroxylamine, 168.1, in which R² is an alkyl, alkenyl, cycloalkyl or

cycloalkenyl group, optionally incorporating a heteroatom O, S or N, or a functional group such as an amide, ester, oxime, sulfoxide or sulfone etc, or an optionally substituted aryl, heteroaryl or aralkyl group, optionally incorporating a heteroatom O, S or N, and X is either a phosphonate group or a group which is subsequently converted into a phosphonate-containing substituent. For example, X is dialkylphosphono, bromo, hydroxy, amino, carboxy and the like. The reaction is conducted between equimolar amounts of the reactants in an aprotic solvent such as pyridine or xylene, or in an alcoholic solvent such as ethanol, optionally in the presence of an acid catalyst, to give the imine or oxime. The preparation of oximes of steroidal 3-ketones is described in *Anal. Bioch.*, 1978, 86, 133 and in *J. Mass. Spectrom.*, 1995, 30, 497. The BMD-protected side-chain compound, 168.2, is then converted into the triol, 168.3, and then to the 21-chloro 17-(2-furoate) product, 168.4, as described herein.



The preparation of hydroxylamine ethers incorporating a phosphonate group is illustrated herein. In this procedure, a phosphonate, 168.5, in which Lv is a leaving group such as bromo or trifluoromethylsulfonyloxy, is reacted with BOC-hydroxylamine, 168.6, (Aldrich) to produce the ether, 168.7. The reaction is conducted between equimolar amounts of the reactants in a polar solvent such as dimethylformamide or tetrahydrofuran, in the presence of a base such as potassium hydroxide or dimethylaminopyridine. Deprotection, for example by treatment with trifluoroacetic acid, then gives the hydroxylamine ether, 168.8.

Example 169

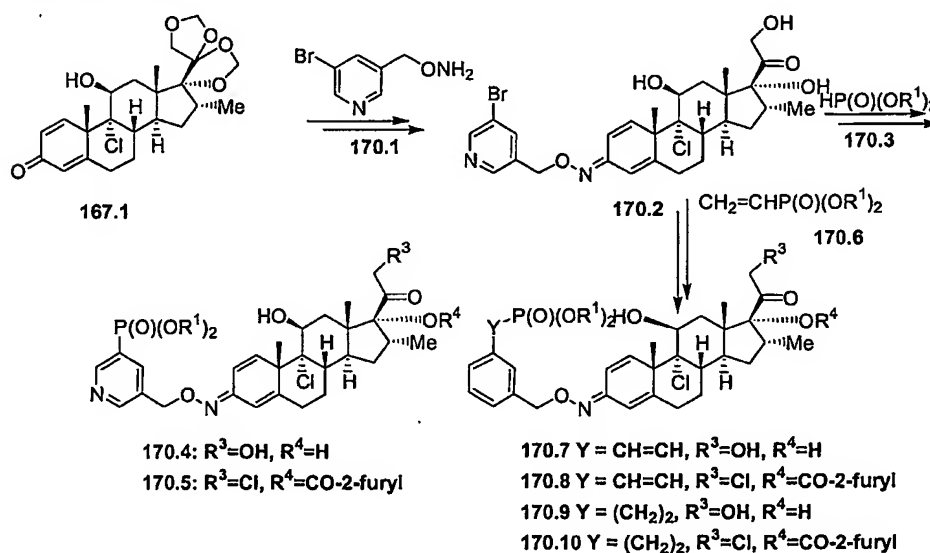


The substrate, 167.1, is reacted with a dialkyl phosphonomethyl hydroxylamine, 169.1, prepared as described above from a dialkyl

trifluoromethylsulfonyloxymethyl phosphonate (*Tet. Lett.*, 1986, 27, 1477) and BOC-hydroxylamine, to afford the oxime, 169.2. Deprotection then affords the triol, 169.3, from which the 21-chloro 17-(2-furoate) compound, 169.4, is prepared, using the procedures described in Example 167. The oxime forming reaction is performed at ambient temperature in ethanol-acetic acid solution between equimolar amounts of the reactants.

Using the above procedures, but employing, in place of the hydroxylamine ether, 169.1, different oxime ethers, 167.1, the corresponding products, 169.4 are obtained.

Example 170



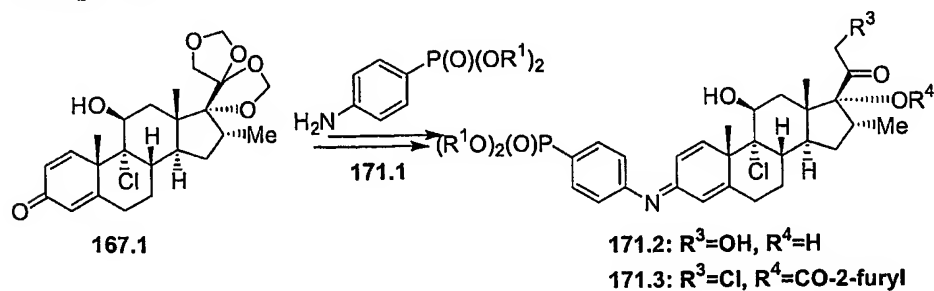
The dienone, 167.1, is reacted, as described above, with O-(5-bromo-3-pyridylmethoxy)hydroxylamine, 170.1, prepared as described above from 5-bromo-3-bromomethylpyridine (EP 511865) and BOC-protected hydroxylamine, 168.6, to give, after deprotection of the side-chain, the oxime, 170.2. The product is then reacted, in the presence of a palladium catalyst, with a dialkyl phosphite, 170.3, to afford the phosphonate, 170.4. The preparation of arylphosphonates by means of a coupling reaction between aryl bromides and dialkyl phosphites is described in *J. Med. Chem.*, 1992, 35, 1371. The reaction is performed at ca. 100°C in an inert solvent such as toluene, in the presence of a base such as triethylamine and a catalytic amount of tetrakis(triphenyl-

phosphine)palladium(0). The 21-hydroxy compound, 170.4, is then converted, as described in Example 167, into the 21-chloro 17-(2-furoate) derivative, 170.5.

Alternatively, the bromo compound, 170.2, is coupled with a dialkyl vinyl phosphonate, 170.6, (Aldrich) to afford the phosphonate, 170.7. The coupling of aryl halides with olefins by means of the Heck reaction is described, for example, in F. A. Carey and R. J. Sundberg, *Advanced Organic Chemistry*, 503ff (Plenum, 2001) and in *Acc. Chem. Res.*, 1979, 12, 146. The aryl bromide and the olefin are coupled in a polar solvent such as dimethylformamide or dioxan, in the presence of a palladium(0) catalyst such as tetrakis(triphenylphosphine)palladium(0) or palladium(II) catalyst such as palladium(II) acetate, and optionally in the presence of a base such as triethylamine or potassium carbonate. Optionally, the double bond present in the product, 170.7, is reduced, for example by reaction with diimide, to produce the saturated analog, 170.9. The reduction of olefinic bonds is described in R. C. Larock, *Comprehensive Organic Transformations*, 6ff (VCH, 1989). The transformation is effected by means of catalytic hydrogenation, for example using a palladium on carbon catalyst and hydrogen or a hydrogen donor, or by the use of diimide or diborane. The products, 170.7 and 170.9, are then converted into the 21-chloro 17-(2-furoate) analogs, 170.8 and 170.10.

Using the above procedures, but employing, in place of the bromopyridylmethoxy reagent, 170.1, different bromo-substituted aryl or heteroaryl alkoxy hydroxylamines, and/or different dialkyl alkenyl phosphonates, the products analogous to the compounds, 170.5, 170.8 and 170.10 are obtained.

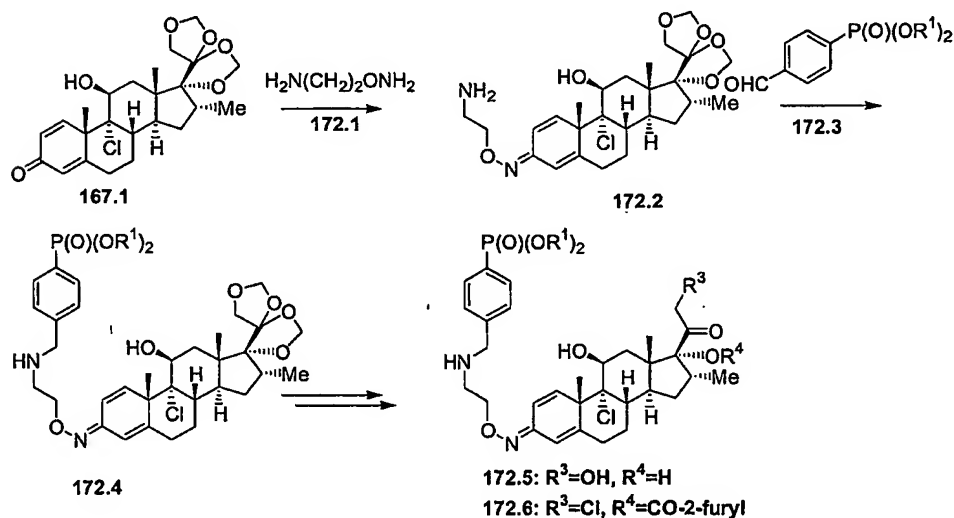
Example 171



The preparation of phosphonates in which the phosphonate is attached by means of an imino group is illustrated herein. In this procedure, the substrate, **167.1**, is reacted with a dialkyl 4-aminophenyl phosphonate, **171.1**, (Epsilon) to give, after deprotection, the imine product, **171.2**. The imine forming reaction is conducted in a hydrocarbon solvent such as toluene or xylene, at reflux temperature, in the presence of a basic catalyst such as sodium methoxide, or an acid catalyst such as p-toluenesulfonic acid, under azeotropic conditions. The product is then converted into the 21-chloro 17-(2-furoate) compound, **171.3**.

Using the above procedures, but employing, in place of the 4-aminophenyl phosphonate, **171.1**, different amino-substituted aryl or heteroaryl phosphonates, products analogous to **171.3** are obtained.

Example 172



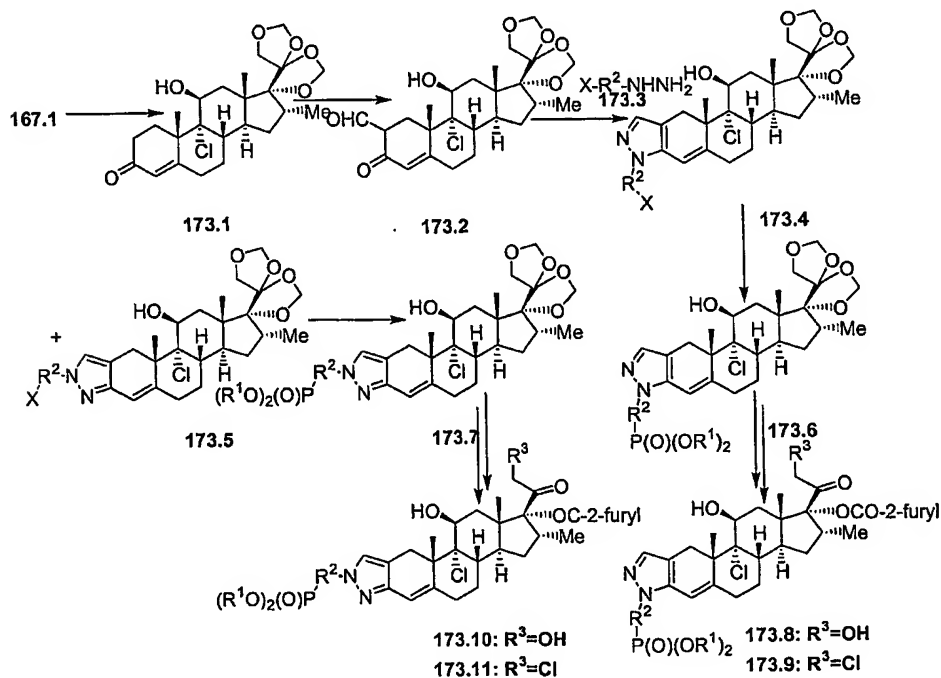
The dienone, **167.1**, is reacted with O-(2-aminoethyl)hydroxylamine, **172.1**, (*Pol. J. Chem.*, 1981, 55, 1163) to yield the oxime, **172.2**. The reaction of steroidal 1,4-dien-3-ones with substituted hydroxylamines is described in *J. Steroid Bioch.*, 1976, 7, 795; the reaction is performed between equimolar amounts of the reactants in a polar organic solvent such as pyridine or methanol, optionally in the presence of acetic acid or sodium acetate. The product is then reacted, in a reductive amination procedure, with a dialkyl 4-formylphenyl phosphonate, **172.3**, (Epsilon) and sodium triacetoxyborohydride, to yield the

amine oxime 172.4. The preparation of amines by means of reductive amination procedures is described, for example, in R. C. Larock, *Comprehensive Organic Transformations*, 421 (VCH), and in F.A. Carey and R. J. Sundberg, *Advanced Organic Chemistry*, Part B, 269 (Plenum, 2001). In this procedure, the amine component and the aldehyde or ketone component are reacted together in the presence of a reducing agent such as, for example, borane, sodium cyanoborohydride, sodium triacetoxyborohydride or diisobutylaluminum hydride, optionally in the presence of a Lewis acid, such as titanium tetraisopropoxide, as described in *J. Org. Chem.*, 1990, 55, 2552.

The amine product, 172.4, is then converted, as described herein, into the 21-chloro 17-(2-furoate) product, 171.6.

Using the above procedures, but employing, in place of the hydroxylamine, 172.3, different amino-substituted hydroxylamines, and/or different formyl-substituted phosphonates, the products analogous to 177.6 are obtained.

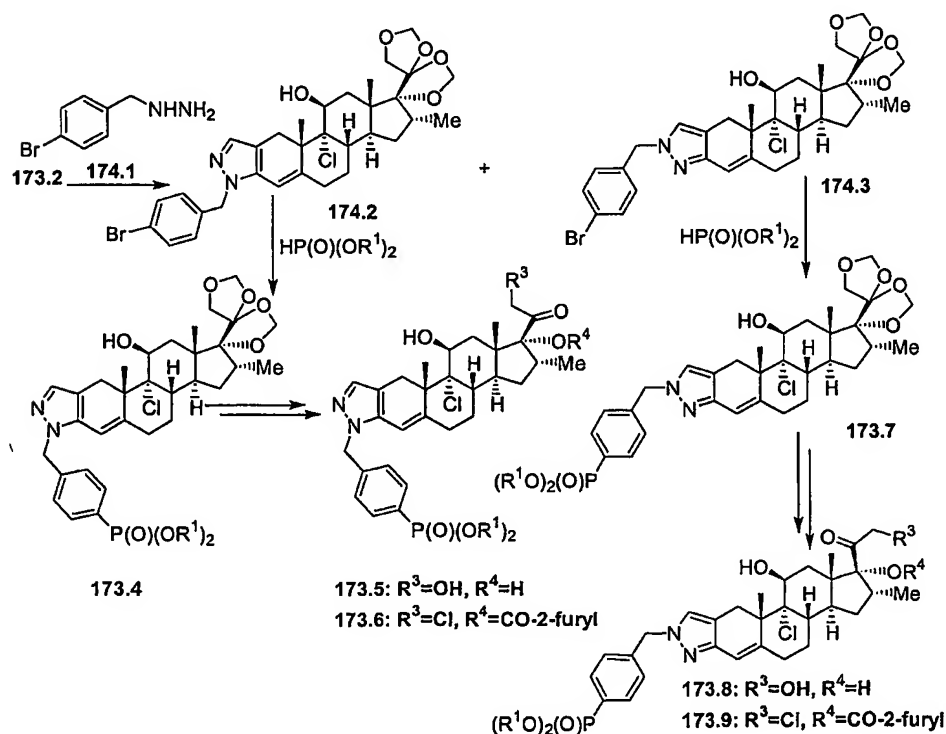
Example 173



The BMD-protected dienone, 167.1, is reduced to afford the 1,2-dihydro product, 173.1. The catalytic hydrogenation reaction is effected by the use of

tris(triphenylphosphine)rhodium (I) chloride, for example as described in *J. Med. Chem.*, **2001**, *44*, 602. The product is then reacted with ethyl formate and a base such as sodium hydride, in an inert solvent such as toluene or dimethylformamide, as described in *J. Am. Chem. Soc.*, **1964**, *86*, 1520, to afford the 2-formyl product, **173.2**. This compound is then reacted with an alkyl, aralkyl, aryl or heteroaryl hydrazine, **173.3**, in which the substituent X is either a phosphonate group or a group which is subsequently transformed into a phosphonate-containing substituent. For example, X is dialkylphosphono, bromo, hydroxy, amino, carboxyl and the like. The reaction yields the isomeric 2'- and 1'-aryl pyrazoles, **173.4** and **173.5**. The pyrazole-forming reaction is performed between equimolar amounts of the reactants in an acidic solvent such as acetic acid, as described in *J. Am. Chem. Soc.*, **1964**, *86*, 1520. The pyrazoles, **173.4** and **173.5**, are then transformed, for example by the procedures described herein, via the BMD-protected intermediates, **173.6** and **173.7**, into the 21-chloro 17-(2-furoate) phosphonates, **173.9** and **173.11**.

Example 174

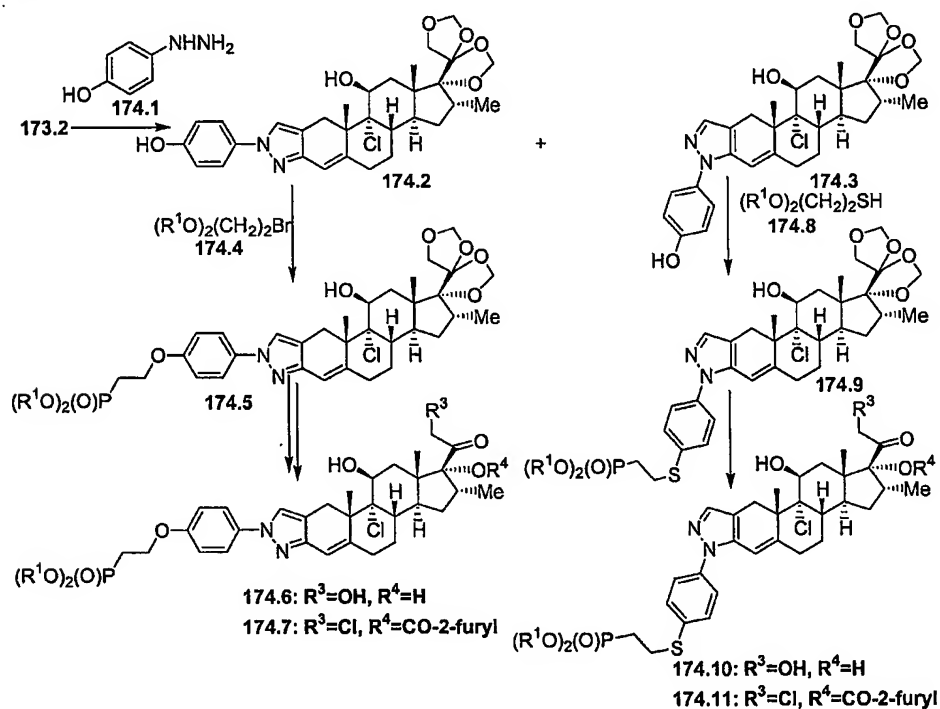


The ketoaldehyde, **173.2**, is reacted, as described above, with 4-bromobenzyl hydrazine, **174.1**, (*Ann.*, 1968, 717, 104) to give the pyrazoles, **174.2** and **174.3**. The 2'-substituted isomer, **174.2**, is then coupled, as described in Example 170, with a dialkyl phosphite, to yield the phosphonate, **173.4**. The BMD protecting group is then removed and the product is converted into the 21-chloro 17-(2-furoate) product, **173.6**.

The isomeric pyrazole, **174.3**, is subjected to the same series of reactions to afford the isomeric product, **173.9**.

Using the above procedures, but employing different bromo-substituted hydrazines, the products analogous to **173.6** and **173.9** are obtained.

Example 174

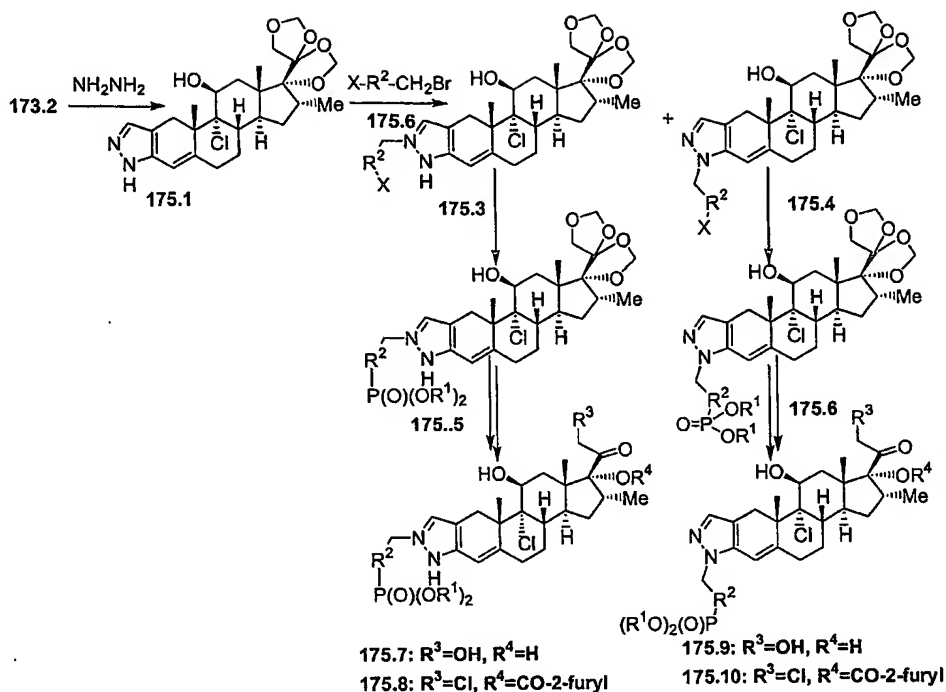


The ketoaldehyde, **173.2**, is reacted, as described above, with 4-hydroxyphenyl hydrazine, **174.1**, (EP 437105) to produce the pyrazoles, **174.2** and **174.3**. The 1'-substituted isomer, **174.2**, is reacted in dimethylformamide at 70°C, with a dialkyl 2-bromoethyl phosphonate, **174.4**, (Aldrich) and potassium carbonate, to give the ether phosphonate, **174.5**. The product is then deprotected to afford

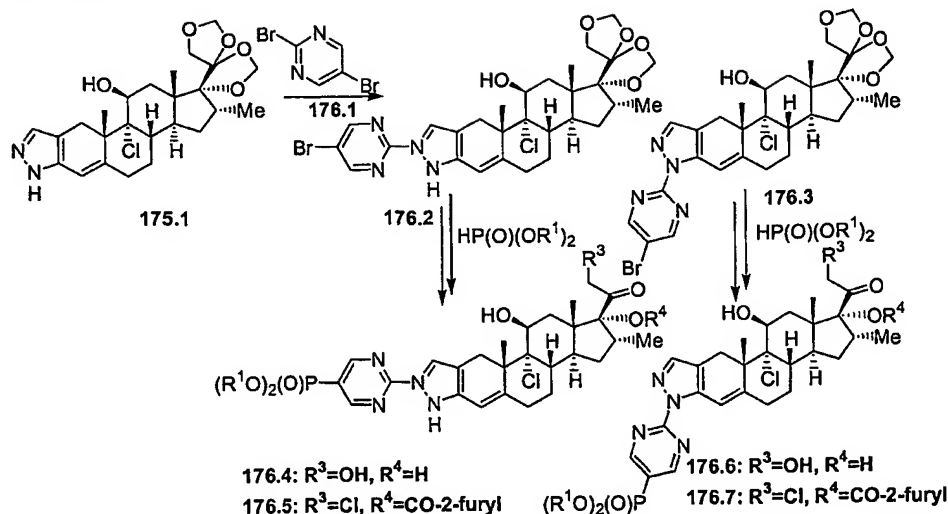
the triol, 174.6, which is converted into the 21-chloro 17-(2-furoate) compound, 174.7.

Alternatively, the 2'-substituted pyrazole, 174.3, is coupled, in a Mitsunobu reaction, with a dialkyl 2-mercaptoethyl phosphonate, 174.8, (*Zh. Obschei. Khim.*, 1973, 43, 2364) to prepare the thioether phosphonate, 174.9, which is deprotected, and the product is converted into the 21-chloro 17-(2-furoate) analog, 174.11. The preparation of aromatic ethers and thioethers by means of the Mitsunobu reaction is described, for example, in R. C. Larock, *Comprehensive Organic Transformations*, 448 (VCH, 1989), and in F.A. Carey and R. J. Sundberg, *Advanced Organic Chemistry*, Part B, 153-4 (Plenum, 2001) and in *Org. React.*, 1992, 42, 335. The phenol and the alcohol or thiol component are reacted together in an aprotic solvent such as, for example, tetrahydrofuran, in the presence of a dialkyl azodicarboxylate and a triarylphosphine, to afford the ether or thioether products. The procedure is also described in *Org. React.*, 1992, 42, 335-656.

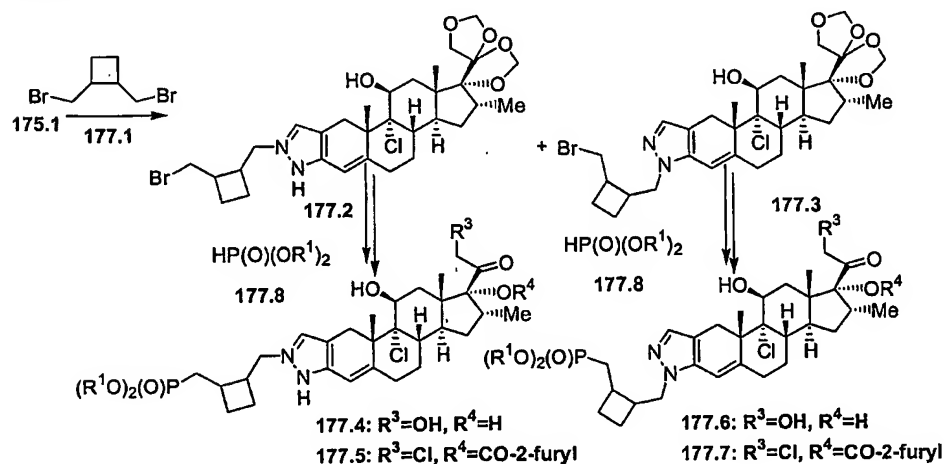
Using the above procedures, but employing, in place of the 4-hydroxyphenyl hydrazine, 174.1, different hydroxy-substituted hydrazines, and/or different dialkyl bromo- or mercapto-substituted phosphonates, the products analogous to the compounds, 174.7 and 174.11 are obtained.

Example 175

The ketoaldehyde, 173.2, is reacted with hydrazine, to afford the pyrazole derivative, 175.1. The reaction of steroidal 2-formyl-3-ketones with hydrazine is described in *J. Am. Chem. Soc.*, 1964, 86, 1520. The reaction is performed in acetic acid at ambient temperature. The pyrazole product is then reacted with a bromomethyl compound, 175.6, in which R^2 and X are as defined above, or a reactive bromoheteroaromatic reagent, to yield the alkylation products, 175.3 and 175.4. The alkylation of substituted pyrazoles is described, for example, in T. L. Gilchrist, *Heterocyclic Chemistry*, 309 (Longman, 1992). The reaction is performed between equimolar amounts of the substrates in a polar solvent such as dimethylformamide or tetrahydrofuran, in the presence of a base such as dimethylaminopyridine, lithium hexamethyldisilazide and the like. The products, 175.3 and 175.4, are, except in cases where X is dialkylphosphono, converted into the phosphonates, 175.5 and 175.6, using the procedures described herein, and deprotection/chlorination/acetylation then affords the 21-chloro 17-(2-furoate) compounds, 175.8 and 175.10.

Example 176

The pyrazole, 175.1, is reacted with 2,5-dibromopyrimidine, 176.1, (*Chem. Lett.*, 1992, 583) to give the pyrazoles, 176.2 and 176.3. The products are then coupled, as described above, with a dialkyl phosphite, to afford after side-chain deprotection and modification, as described above, the 21-chloro 17-(2-furoates), 176.5 and 176.7.

Example 177

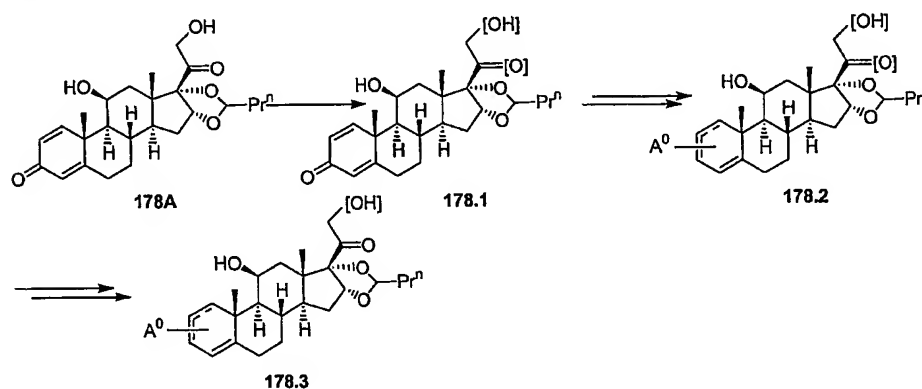
The pyrazole, 175.1, is reacted in tetrahydrofuran solution, with 1,2-bis(bromomethyl)cyclobutane, 177.1 (*J. Org. Chem.*, 1981, 46, 3530) and potassium hexamethyl disilazide, to give the alkylation products, 177.1 and 177.2. The 1'-substituted isomer, 177.2, is then reacted, in an Arbuzov reaction, with a trialkyl phosphite to yield, after deprotection and side-chain modification,

the 21-chloro 17-(2-furoate), **177.5**. The Arbuzov reaction is described in *Handb. Organophosphorus Chem.*, **1992**, 115. In this procedure, in which a bromo substituent is converted into the corresponding phosphonate, the substrate is heated at from about 60°C to about 160°C with a five to fifty-fold molar excess of a trialkyl phosphite, to effect the transformation.

The 2'-substituted pyrazole, **177.3**, is subjected to the same series of reaction to give the amine phosphonate, **177.7**.

Using the above procedures, but employing different dibromides, the products analogous to **177.5** and **177.7** are obtained.

Example 178



A protection-deprotection sequence in which the 20-ketone group and/or the 21-hydroxyl group of Budesonide, **178A**, are protected to afford the derivative, **178.1**. The ketone is protected, for example, by conversion to the cyclic ethylene ketal, by reaction in toluene solution at reflux temperature with ethylene glycol and an acid catalyst, as described in *J. Am. Chem. Soc.*, **1955**, 77, 1904. Deprotection is effected by reaction with pyridinium tosylate in aqueous acetone, as described in *J. Chem. Soc. Chem. Comm.*, **1987**, 1351.

Alternatively, the 20-ketone is protected by conversion to the N, N-dimethylhydrazone. The dimethyl hydrazone is prepared by the reaction of the ketone, **178A**, with N,N-dimethylhydrazine in ethanol-acetic acid, as described in *Org. Syn.*, **1970**, 50, 102. The group is removed by treatment with sodium acetate and acetic acid in aqueous tetrahydrofuran, as described in *J. Am. Chem. Soc.*, **1979**, 101, 5841.

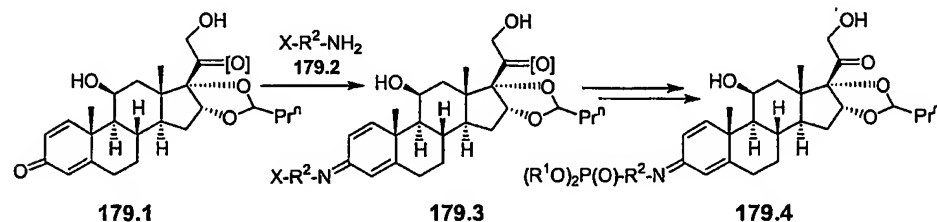
Alternatively, the 20-ketone is protected as the diethylamine adduct. In this procedure, the substrate, 178A, is reacted with titanium tetrakis(diethylamide), as described in *J. Chem. Soc., Chem. Comm.*, 1983, 406, to afford the adduct. The ketone is deprotected by reaction with water in an aqueous organic solvent.

The 21-hydroxyl group is protected, for example, by conversion to the acetate ester, by reaction with one molar equivalent of acetyl chloride in dichloromethane/pyridine. The 21-acetoxy group is removed by reaction with one molar equivalent of lithium hydroxide in aqueous dimethoxyethane.

Alternatively, the 21-hydroxyl group is protected by conversion to the tert. butyl dimethylsilyl ether, by reaction in dimethylformamide solution with one molar equivalent of tert. butylchlorodimethylsilane and imidazole, as described in *J. Am. Chem. Soc.*, 1972, 94, 6190. The silyl ether is removed by reaction with tetrabutylammonium fluoride in tetrahydrofuran solution, as described in *J. Am. Chem. Soc.*, 1972, 94, 6190.

The protected compound, 178.1, is then converted into the phosphonate-containing analog, 178.2, using the procedures described below, and the protecting group or groups are then removed, as described above, to give the phosphonate, 178.3.

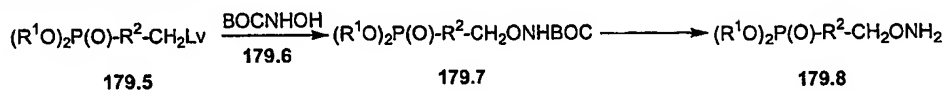
Example 179



The ketone-protected derivative, 179.1, is reacted with an amine or hydroxylamine, 179.2, in which R^2 is an alkyl, alkenyl, cycloalkyl or cycloalkenyl group, optionally incorporating a heteroatom O, S or N, or a functional group such as an amide, ester, oxime, sulfoxide or sulfone etc, or an optionally substituted aryl, heteroaryl or aralkyl group, optionally incorporating a heteroatom O, S or N, and X is either a phosphonate group or a group which is subsequently converted into a phosphonate-containing substituent. For example, X is dialkylphosphono, bromo, hydroxy, amino, carboxy and the like. The

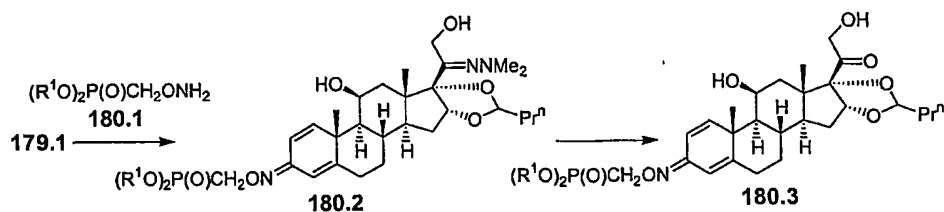
reaction is conducted between equimolar amounts of the reactants in an aprotic solvent such as pyridine or xylene, or in an alcoholic solvent such as ethanol, optionally in the presence of an acid catalyst, to give the imine or oxime, 179.3. The preparation of oximes of steroidal 3-ketones is described in *Anal. Bioch.*, 1978, 86, 133 and in *J. Mass. Spectrom.*, 1995, 30, 497. The protecting group is then removed, as described herein, to afford the 20-keto phosphonate product, 179.4.

Example 179A



The preparation of hydroxylamine ethers incorporating a phosphonate group is also illustrated. In this procedure, a phosphonate, 179.5, in which Lv is a leaving group such as bromo or trifluoromethylsulfonyloxy, is reacted with BOC-hydroxylamine, 179.6, (Aldrich) to produce the ether, 179.7. The reaction is conducted between equimolar amounts of the reactants in a polar solvent such as dimethylformamide or tetrahydrofuran, in the presence of a base such as potassium hydroxide or dimethylaminopyridine. Deprotection, for example by treatment with trifluoroacetic acid, then gives the hydroxylamine ether, 179.8. The above procedure is also employed for the preparation of substituted hydroxylamines which are precursors to phosphonates.

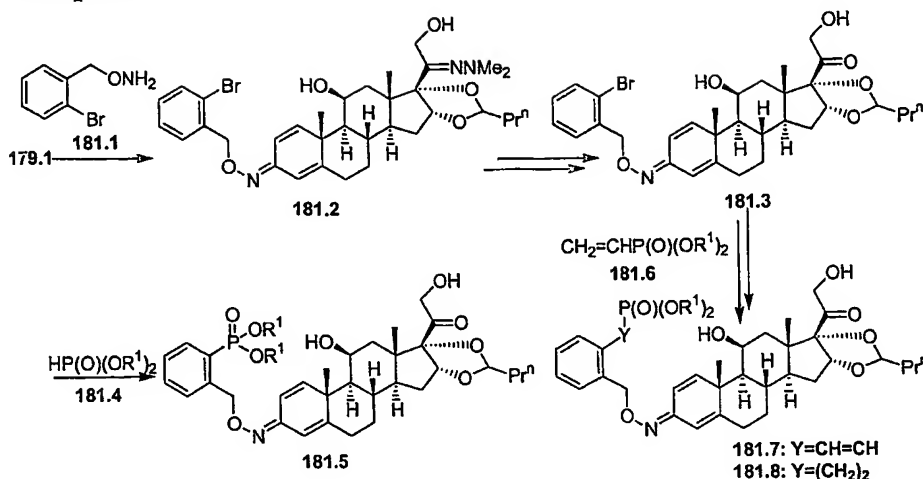
Example 180



The substrate, 179.1, in which the 20-ketone is protected as the dimethyl hydrazone derivative, is reacted with a dialkyl phosphonomethyl hydroxylamine, 180.1, prepared as described above from a dialkyl trifluoromethylsulfonyloxymethyl phosphonate (*Tet. Lett.*, 1986, 27, 1477) and BOC-hydroxylamine, to afford the oxime, 180.2. Deprotection, as described in Example 179, affords the

20-keto phosphonate, **180.3**. The oxime forming reaction is performed at ambient temperature in ethanol-acetic acid solution between equimolar amounts of the reactants.

Using the above procedures, but employing, in place of the hydroxylamine ether, **180.1**, different oxime ethers, **179.2**, the corresponding products, **179.4** are obtained.

Example 181

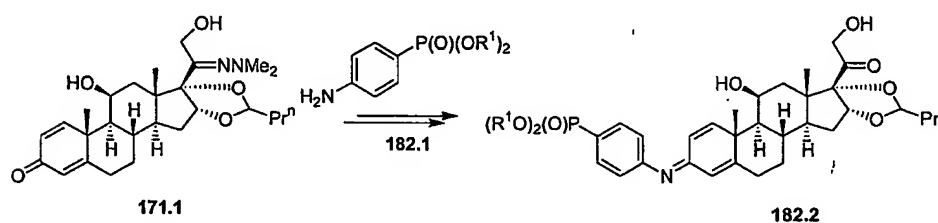
The preparation of compounds, **179.1**, in which the phosphonate group is attached by means of a benzyloxy oxime group is illustrated above. In this procedure, the dienone, **179.1**, in which the 20-ketone is protected as the dimethyl hydrazone, is reacted, as described above, with O-(2-bromobenzyl)hydroxylamine, **181.1**, prepared as described above from 2-bromobenzyl bromide and BOC-protected hydroxylamine, **179.6**, to give the oxime, **181.2**. The protecting group is then removed to yield the 20-keto product, **181.3**. The latter product is then reacted, in the presence of a palladium catalyst, with a dialkyl phosphite, **181.4**, to afford the phosphonate, **181.5**. The preparation of arylphosphonates by means of a coupling reaction between aryl bromides and dialkyl phosphites is described in *J. Med. Chem.*, **1992**, *35*, 1371. The reaction is performed at ca. 100° in an inert solvent such as toluene, in the presence of a base such as triethylamine and a catalytic amount of tetrakis(triphenylphosphine)palladium(0).

Alternatively, the bromo compound, **181.3**, is coupled with a dialkyl vinylphosphonate, **181.6**, (Aldrich) to afford the phosphonate, **181.7**. The coupling of aryl halides with olefins by means of the Heck reaction is described, for example, in *Advanced Organic Chemistry*, by F. A. Carey and R. J. Sundberg, Plenum, **2001**, p. 503ff and in *Acc. Chem. Res.*, **1979**, *12*, 146. The aryl bromide and the olefin are coupled in a polar solvent such as dimethylformamide or dioxan, in the presence of a palladium(0) catalyst such as tetrakis(triphenylphosphine)palladium(0) or palladium(II) catalyst such as

palladium(II) acetate, and optionally in the presence of a base such as triethylamine or potassium carbonate. Optionally, the styrenoid double bond present in the product, **181.7**, is reduced, for example by reaction with diimide, to produce the saturated analog, **181.8**. The reduction of olefinic bonds is described in *Comprehensive Organic Transformations*, by R. C. Larock, VCH, 1989, p. 6ff. The transformation is effected by means of catalytic hydrogenation, for example using a palladium on carbon catalyst and hydrogen or a hydrogen donor, or by the use of diimide or diborane.

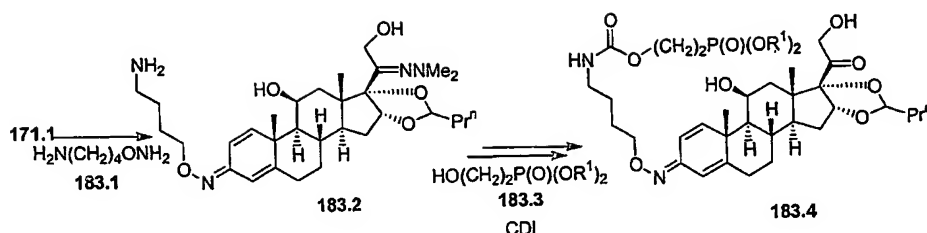
Using the above procedures, but employing, in place of the bromobenzyl reagent, **181.1**, different bromo-substituted aryl or heteroaryl alkoxy hydroxylamines, and/or different dialkyl alkenyl phosphonates, the products analogous to the compounds, **181.5**, **181.7** and **181.8** are obtained.

Example 182



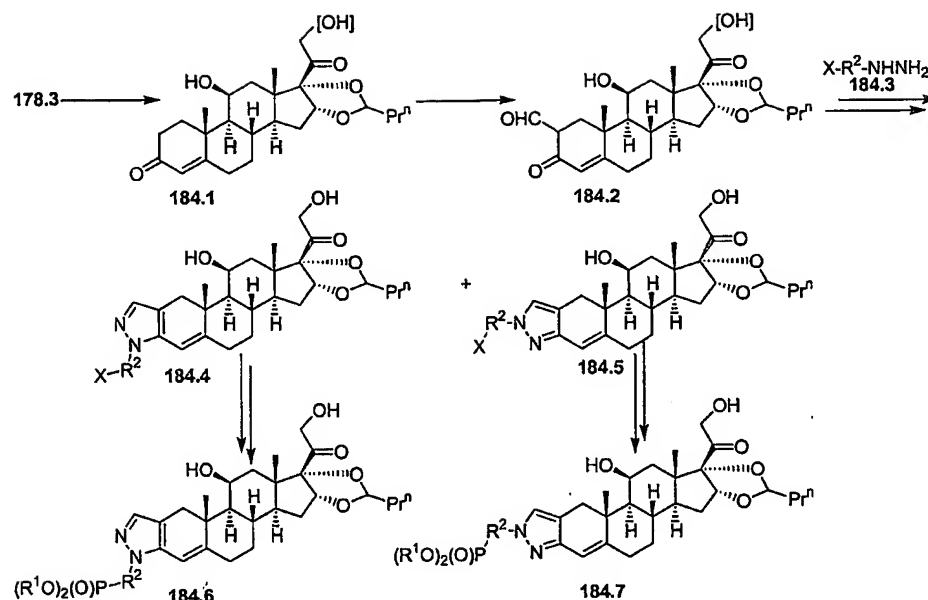
The substrate, **171.1**, in which the 20-ketone is protected as the dimethylhydrazone, is reacted with a dialkyl 4-aminophenyl phosphonate, **182.1**, (Epsilon), to give, after deprotection, the imine product, **182.2**. The imine forming reaction is conducted in a hydrocarbon solvent such as toluene or xylene, at reflux temperature, in the presence of a basic catalyst such as sodium methoxide, or an acid catalyst such as p-toluenesulfonic acid, under azeotropic conditions.

Using the above procedures, but employing, in place of the 4-amino-phenyl phosphonate, **182.1** different amino-substituted aryl or heteroaryl phosphonates, products analogous to **182.2** are obtained.

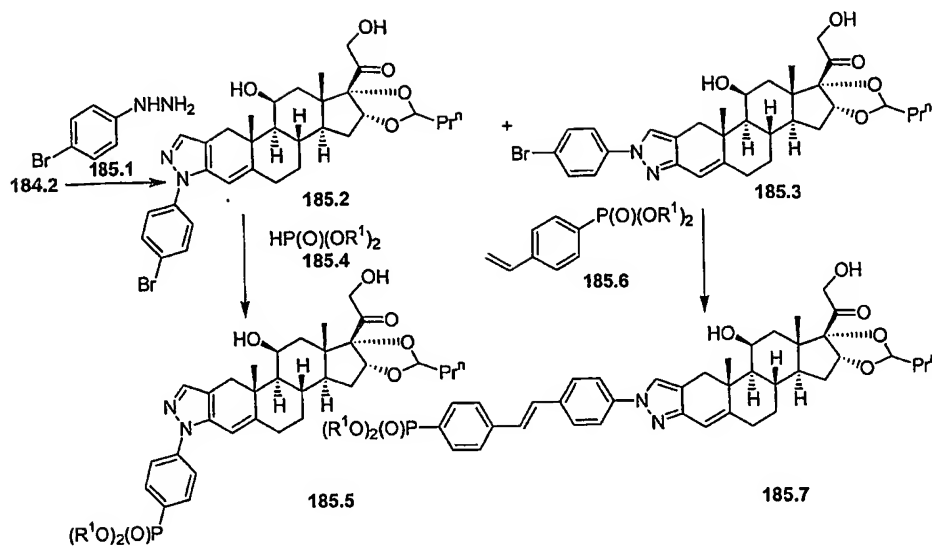
Example 183

The preparation of phosphonates in which the phosphonate is attached by means of an oximino group and a carbamate linkage is illustrated herein. In this procedure, the dienone, 171.1, in which the 20-ketone is protected as the dimethylhydrazone, is reacted with 4-aminobutyl hydroxylamine, 183.1, (*Pol. J. Chem.*, 1981, 55, 1163) to yield the oxime, 183.2. (The reaction of steroidal 1,4-dien-3-ones with hydroxylamines is described in *J. Steroid Bioch.*, 1976, 7, 795.) The reaction is performed between equimolar amounts of the reactants in a polar organic solvent such as pyridine or methanol, optionally in the presence of acetic acid or sodium acetate. The product, 183.2, is then coupled with a dialkyl 2-hydroxyethyl phosphonate, 183.3, (Epsilon) and carbonyl diimidazole (CDI), to yield, after deprotection, the carbamate oxime, 183.4. The preparation of carbamates is described in *Comprehensive Organic Functional Group Transformations*, A. R. Katritzky, ed., Pergamon, 1995, Vol. 6, p 416ff, and in *Organic Functional Group Preparations*, by S. R. Sandler and W. Karo, Academic Press, 1986, p. 260ff. In the procedure, the amine is reacted in an inert aprotic solvent such as dichloromethane or tetrahydrofuran, with phosgene or a functional equivalent thereof, such as carbonyl diimidazole, triphosgene, pentafluorophenyl carbonate and the like, to afford the corresponding activated acylamine. The latter compound is then reacted with an alcohol to yield the carbamate.

Using the above procedures, but employing, in place of the amino-substituted hydrazine, 183.1, different amino-substituted hydrazines, and/or different hydroxy-substituted phosphonates, the products analogous to 183.4 are obtained.

Example 184

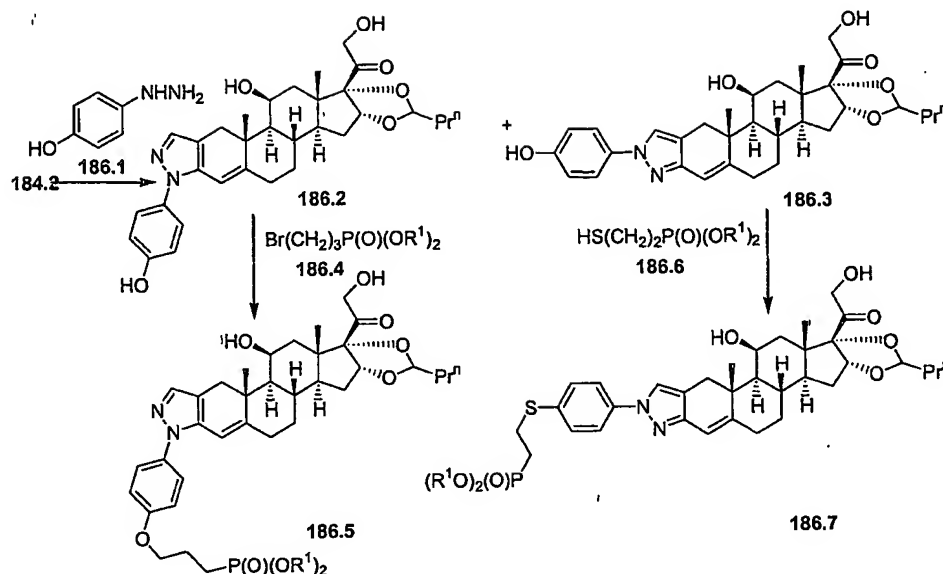
The dienone, 178.3, in which the 21-hydroxyl group is protected as described in Example 178 is reduced to afford the 1,2-dihydro product, 184.1. The catalytic hydrogenation reaction is effected by the use of tris(triphenylphosphine)rhodium (I) chloride, for example as described in *J. Med. Chem.*, 2001, 44, 602. The product is then reacted with ethyl formate and a base such as sodium hydride, in an inert solvent such as toluene or dimethylformamide, as described in *J. Am. Chem. Soc.*, 1964, 86, 1520, to afford the 2-formyl product, 184.2. This compound is then reacted with an alkyl, aralkyl, aryl or heteroaryl hydrazine, 184.3, in which the substituent X is either a phosphonate group or a group which is subsequently transformed into a phosphonate-containing substituent. For example, X is dialkylphosphono, bromo, hydroxy, amino, carboxyl and the like. The reaction yields, after deprotection of the 21-hydroxyl group, the isomeric 2'- and 1'-aryl pyrazoles, 184.4 and 184.5. The pyrazole-forming reaction is performed between equimolar amounts of the reactants in an acidic solvent such as acetic acid, as described in *J. Am. Chem. Soc.*, 1964, 86, 1520. The pyrazoles, 184.4 and 184.5, are then transformed, for example by the procedures described in Examples 180 and 181, into the phosphonates, 184.6 and 184.7.

Example 185

The ketoaldehyde, **184.2**, is reacted, as described above, with 4-bromophenyl hydrazine, **185.1**, (*J. Organomet. Chem.*, **1999**, *62*, 581) to give the pyrazoles, **185.2** and **185.3**. The 2'-substituted isomer **185.2** is then reacted, as described above, with a dialkyl phosphate, **184.4**, to give the phosphonate, **185.5**.

The isomeric pyrazole, **185.3**, is reacted in a Heck reaction, as described above, with one molar equivalent of a dialkyl 4-vinylphenyl phosphonate, **185.6**, (*Macromolecules*, **1998**, *31*, 2918) to yield the phosphonate, **185.7**.

Using the above procedures, but employing different bromo-substituted hydrazines, and/or different alkenyl-substituted phosphonates, the products analogous to **185.5** and **185.7** are obtained.

Example 186

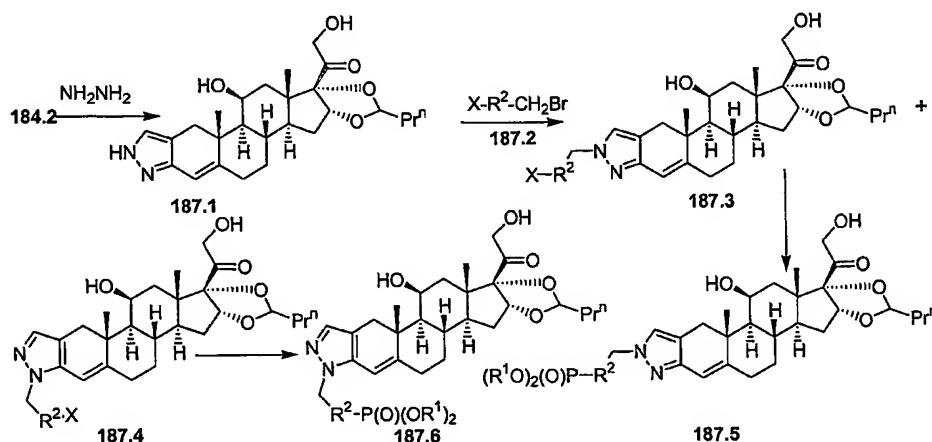
The ketoaldehyde, **184.2**, is reacted, as described above, with 4-hydroxyphenyl hydrazine, **186.1**, (EP 437105) to produce the pyrazoles, **186.2** and **186.3**. The 2'-substituted isomer, **186.2**, is then reacted in dimethylformamide solution at 70° with one molar equivalent of a dialkyl bromopropyl phosphonate, **186.4**, (*J. Amer. Chem. Soc.*, 2000, 122, 1554) and cesium carbonate, to give the ether phosphonate, **186.5**.

Alternatively, the 1'-substituted pyrazole, **186.3**, is coupled in a Mitsunobu reaction, with a dialkyl 2-mercaptoethyl phosphonate, **186.6**, (*Zh. Obshchei. Khim.*, 1973, 43, 2364) to prepare the thioether phosphonate, **186.7**. The preparation of aromatic ethers and thioethers by means of the Mitsunobu reaction is described, for example, in *Comprehensive Organic Transformations*, by R. C. Larock, VCH, 1989, p. 448, and in *Advanced Organic Chemistry*, Part B, by F.A. Carey and R. J. Sundberg, Plenum, 2001, p. 153-4 and in *Org. React.*, 1992, 42, 335. The phenol and the alcohol or thiol component are reacted together in an aprotic solvent such as, for example, tetrahydrofuran, in the presence of a dialkyl azodicarboxylate and a triarylphosphine, to afford the ether or thioether products. The procedure is also described in *Org. React.*, 1992, 42, 335-656.

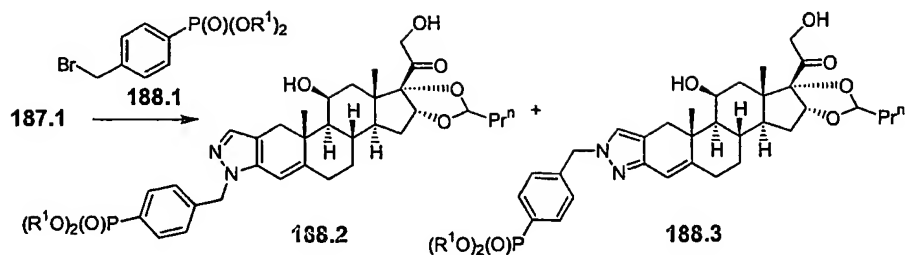
Using the above procedures, but employing, in place of the hydroxylphenyl hydrazine, **186.1**, different hydroxyaryl hydrazines, and/or different

dialkyl bromo- or mercapto-substituted phosphonates, the products analogous to the compounds, 186.5 and 186.7 are obtained.

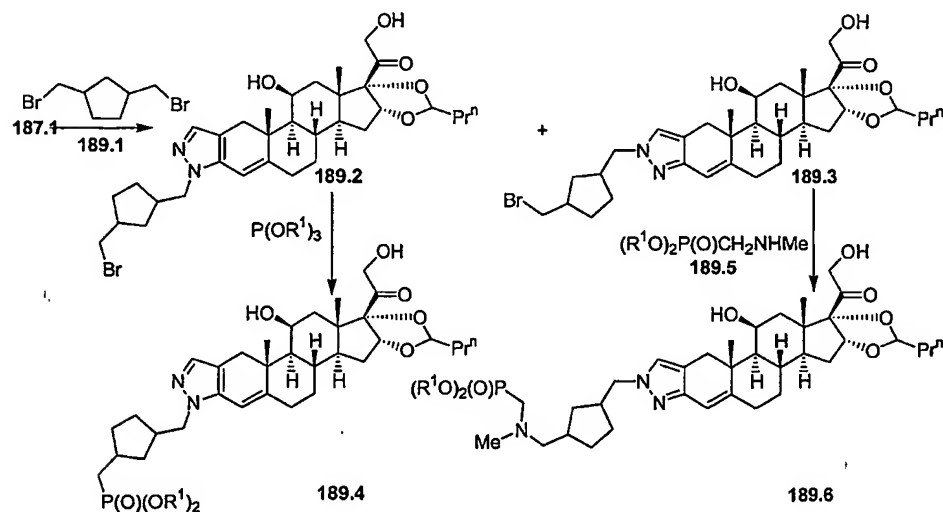
Example 187



The ketoaldehyde, 184.2, is reacted with hydrazine to afford the pyrazole derivative, 187.1. The reaction of steroidal 2-formyl-3-ketones with hydrazine is described in *J. Am. Chem. Soc.*, 1964, 86, 1520. The reaction is performed in acetic acid at ambient temperature. The pyrazole product is then reacted with a bromomethyl compound, 187.2, in which R^2 and X are as defined above, or a reactive bromoheteroaromatic reagent, to yield the alkylation products, 187.3 and 187.4. The alkylation of substituted pyrazoles is described, for example, in *Heterocyclic Chemistry*, by T. L. Gilchrist, Longman, 1992, p. 309. The reaction is performed between equimolar amounts of the substrates in a polar solvent such as dimethylformamide or tetrahydrofuran, in the presence of a base such as dimethylaminopyridine, lithium hexamethyldisilazide and the like. The products, 187.3 and 187.4, are, except in cases where X is dialkylphosphono, converted into the phosphonates, 187.5 and 187.6, using the procedures described herein.

Example 188

The pyrazole, 187.1, is reacted in dimethylformamide solution at 70° with one molar equivalent of a dialkyl 4-bromomethylphenyl phosphonate, 188.1, (*Tet.*, 1998, 54, 9341) and lithium hexamethyl disilazide, to give the pyrazoles, 188.2 and 188.3. Using the above procedures, but employing different bromomethyl-substituted phosphonates, the products analogous to 188.2 and 188.3 are obtained.

Example 189

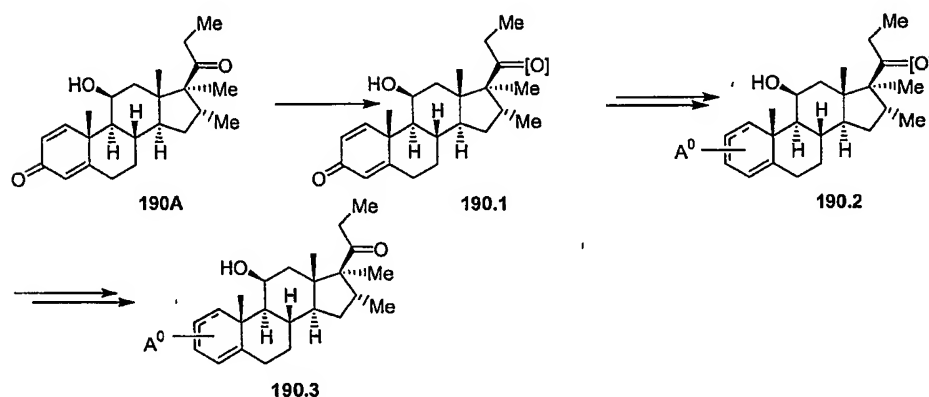
The pyrazole, 187.1, is reacted in tetrahydrofuran solution with 1,3-bis(bromomethyl)cyclopentane, 189.1, (*Bull. Soc. Chim. Fr.*, 1975, 1295) and sodium hydride, to give the alkylation products, 189.2 and 189.3. The 2'-substituted isomer, 189.2, is then reacted, in a Arbuzov reaction, with a trialkyl phosphite to yield the phosphonate, 189.4. The Arbuzov reaction is described in *Handb. Organophosphorus Chem.*, 1992, 115. In this procedure, in which a bromo substituent is converted into the corresponding phosphonate, the substrate

is heated at from about 60° to about 160° with a five to fifty-fold molar excess of a trialkyl phosphite, to effect the transformation.

The 2'-substituted pyrazole, **189.3**, is reacted at 70° in dimethyl-formamide solution with one molar equivalent of a dialkyl methylaminomethyl phosphonate, **189.5**, and cesium carbonate, to give the amine phosphonate, **189.6**.

Using the above procedures, but employing different dihalides, and/or different amino-substituted phosphonates, the products analogous to **189.4** and **189.6** are obtained.

Example 190



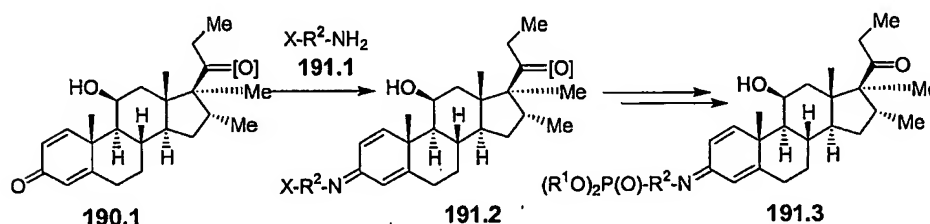
A protection-deprotection sequence in which the 20-ketone group of Rimexolone, **190A**, is protected to afford the derivative, **190.1**. The ketone is protected, for example, by conversion to the cyclic ethylene ketal, by reaction in toluene solution at reflux temperature with ethylene glycol and an acid catalyst, as described in *J. Am. Chem. Soc.*, **1955**, 77, 1904. Deprotection is effected by reaction with pyridinium tosylate in aqueous acetone, as described in *J. Chem. Soc., Chem. Comm.*, **1987**, 1351.

Alternatively, the 20-ketone is protected by conversion to the N, N-dimethylhydrazone. The dimethyl hydrazone is prepared by the reaction of the ketone, **190A**, with N,N-dimethylhydrazine in ethanol-acetic acid, as described in *Org. Syn.*, **1970**, 50, 102. The group is removed by treatment with sodium acetate and acetic acid in aqueous tetrahydrofuran, as described in *J. Am. Chem. Soc.*, **1979**, 101, 5841.

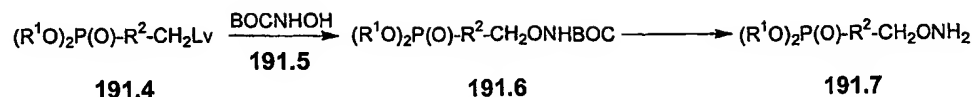
Alternatively, the 20-ketone is protected as the diethylamine adduct. In this procedure, the substrate, **190.1**, is reacted with titanium tetrakis-(diethylamide), as described in *J. Chem. Soc., Chem. Comm.*, **1983**, 406, to afford the adduct. The ketone is deprotected by reaction with water in an aqueous organic solvent.

The protected compound, **190.2**, is then converted into the phosphonate-containing analog, **190.3**, using the procedures described below, and the protecting group or groups are then removed, as described above, to give the phosphonate, **190.3**.

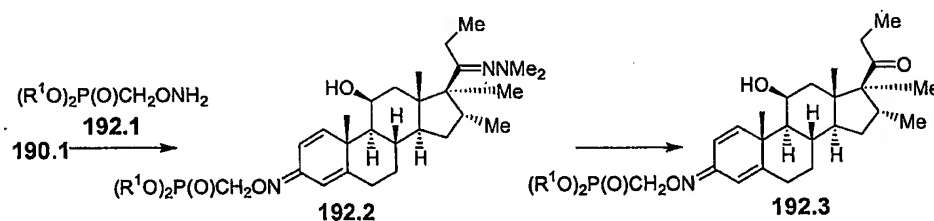
Example 191



The ketone-protected derivative, **190.1**, is reacted with an amine or hydroxylamine, **191.1**, in which R^2 is an alkyl, alkenyl, cycloalkyl or cycloalkenyl group, optionally incorporating a heteroatom O, S or N, or a functional group such as an amide, ester, oxime, sulfoxide or sulfone etc, or an optionally substituted aryl, heteroaryl or aralkyl group, optionally incorporating a heteroatom O, S or N, and X is either a phosphonate group or a group which is subsequently converted into a phosphonate-containing substituent. For example, X is dialkylphosphono, bromo, hydroxy, amino, carboxy and the like. The reaction is conducted between equimolar amounts of the reactants in an aprotic solvent such as pyridine or xylene, or in an alcoholic solvent such as ethanol, optionally in the presence of an acid catalyst, to give the imine or oxime, **191.2**. The preparation of oximes of steroidal 3-ketones is described in *Anal. Bioch.*, **1978**, 86, 133 and in *J. Mass. Spectrom.*, **1995**, 30, 497. The protecting group is then removed, as described in Example 190, to afford the 20-keto phosphonate product, **191.3**.

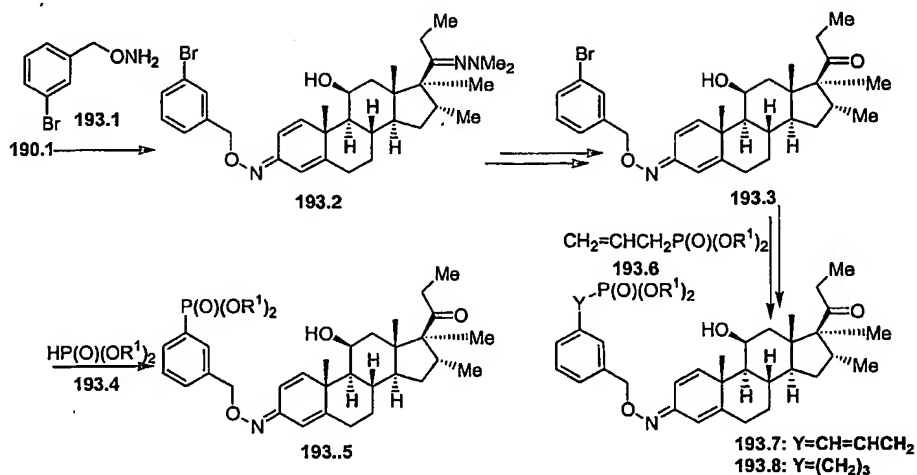
Example 191A

The preparation of hydroxylamine ethers incorporating a phosphonate group is also illustrated. In this procedure, a phosphonate, **191.4**, in which Lv is a leaving group such as bromo or trifluoromethylsulfonyloxy, is reacted with BOC-hydroxylamine, **191.5**, (Aldrich) to produce the ether, **191.6**. The reaction is conducted between equimolar amounts of the reactants in a polar solvent such as dimethylformamide or tetrahydrofuran, in the presence of a base such as potassium hydroxide or dimethylaminopyridine. Deprotection, for example by treatment with trifluoroacetic acid, then gives the hydroxylamine ether, **191.7**. The above procedure is also employed for the preparation of substituted hydroxylamines which are precursors to phosphonates.

Example 192

The substrate, **190.1**, in which the 20-ketone is protected as the dimethyl hydrazone derivative, is reacted with a dialkyl phosphonomethyl hydroxylamine, **192.1**, prepared as described above from a dialkyl trifluoromethyl-sulfonyloxymethyl phosphonate (*Tet. Lett.*, **1986**, 27, 1477) and BOC-hydroxylamine, to afford the oxime, **192.2**. Deprotection, as described in Example **191**, then affords the 20-keto phosphonate, **192.3**. The oxime forming reaction is performed at ambient temperature in ethanol-acetic acid solution between equimolar amounts of the reactants.

Using the above procedures, but employing, in place of the hydroxylamine ether, **192.1**, different oxime ethers, **191.1**, the corresponding products, **191.3** are obtained.

Example 193

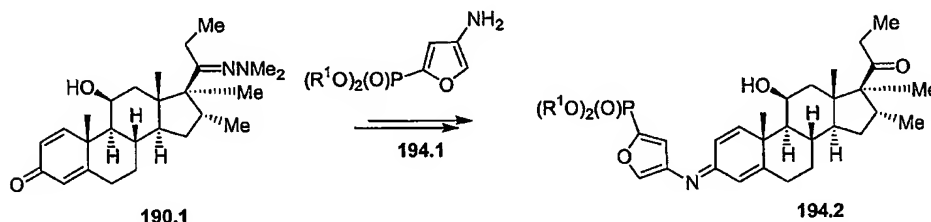
The dienone, 190.1, in which the 20-ketone is protected as the dimethyl hydrazone, is reacted, as described above, with O-(3-bromobenzyl)hydroxylamine, 193.1, prepared as described above from 3-bromobenzyl bromide and BOC-protected hydroxylamine, 191.5, to give the oxime, 193.2. The protecting group is then removed to yield the 20-keto product 193.3. The latter product is then reacted, in the presence of a palladium catalyst, with a dialkyl phosphite, 193.4, to afford the phosphonate, 193.5. The preparation of arylphosphonates by means of a coupling reaction between aryl bromides and dialkyl phosphites is described in *J. Med. Chem.*, **1992**, 35, 1371. The reaction is performed at ca. 100°C in an inert solvent such as toluene, in the presence of a base such as triethylamine and a catalytic amount of tetrakis(triphenylphosphine)-palladium(0).

Alternatively, the bromo compound, 193.3, is coupled with a dialkyl propenylphosphonate, 193.6, (Aldrich) to afford the phosphonate, 193.7. The coupling of aryl halides with olefins by means of the Heck reaction is described, for example, in F. A. Carey and R. J. Sundberg, *Advanced Organic Chemistry*, 503ff (Plenum, 2001) and in *Acc. Chem. Res.*, **1979**, 12, 146. The aryl bromide and the olefin are coupled in a polar solvent such as dimethylformamide or dioxan, in the presence of a palladium(0) catalyst such as tetrakis(triphenylphosphine)palladium(0) or palladium(II) catalyst such as palladium(II) acetate, and optionally in the presence of a base such as triethylamine or potassium carbonate. Optionally, the styrenoid double bond present in the product, 193.7,

is reduced, for example by reaction with diimide, to produce the saturated analog, 193.8. The reduction of olefinic bonds is described in R. C. Larock, *Comprehensive Organic Transformations*, 6ff (VCH, 1989). The transformation is effected by means of catalytic hydrogenation, for example using a palladium on carbon catalyst and hydrogen or a hydrogen donor, or by the use of diimide or diborane.

Using the above procedures, but employing, in place of the bromobenzyl reagent, 193.1, different bromo-substituted aryl or heteroaryl alkoxy hydroxylamines, and/or different dialkyl alkenyl phosphonates, the products analogous to the compounds, 193.5, 193.7 and 193.8 are obtained.

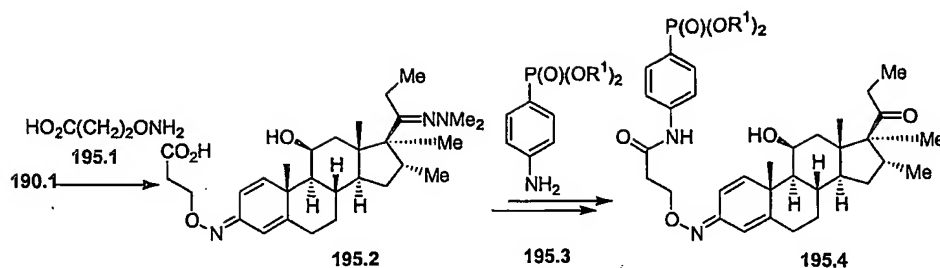
Example 194



The substrate, 190.1, in which the 20-ketone is protected as the dimethylhydrazone, is reacted with a dialkyl 4-amino-2-furyl phosphonate, 194.1, prepared by the palladium catalyzed coupling reaction, as described above, between 4-amino-2-bromofuran (*Tet.*, 1987, 43, 3295) and a dialkyl phosphite, to give, after deprotection, the imine product, 194.2. The imine forming reaction is conducted in a hydrocarbon solvent such as toluene or xylene, at reflux temperature, in the presence of a basic catalyst such as sodium methoxide, or an acid catalyst such as p-toluenesulfonic acid, under azeotropic conditions.

Using the above procedures, but employing, in place of the 4-aminofuryl phosphonate, 194.1, different amino-substituted aryl or heteroaryl phosphonates, products analogous to 194.2 are obtained.

Example 195



The dienone, **190.1**, in which the 20-ketone is protected as the dimethylhydrazone, is reacted with 2-carboxyethyl hydroxylamine, **195.1**, (*J. Med. Chem.*, **1990**, 33, 1423) to yield the oxime, **195.2**. The reaction of steroidal 1,4-dien-3-ones with hydroxylamines is described in *J. Steroid Bioch.*, **1976**, 7, 795; the reaction is performed between equimolar amounts of the reactants in a polar organic solvent such as pyridine or methanol, optionally in the presence of acetic acid or sodium acetate. The product, **195.2**, is then coupled with a dialkyl 4-aminophenyl phosphonate, **195.3**, (Epsilon) and dicyclohexyl carbodiimide, to yield, after deprotection, the amide oxime, **195.4**. The preparation of amides from carboxylic acids and derivatives is described, for example, in S.R. Sandler and W. Karo, *Organic Functional Group Preparations*, 274 (Academic Press, 1968), and R. C. Larock, *Comprehensive Organic Transformations*, 972ff (VCH, 1989). The carboxylic acid is reacted with the amine in the presence of an activating agent, such as, for example, dicyclohexylcarbodiimide or diisopropylcarbodiimide, optionally in the presence of, for example, hydroxybenztriazole, N-hydroxysuccinimide or N-hydroxypyridone, in a non-protic solvent such as, for example, pyridine, DMF or dichloromethane, to afford the amide.

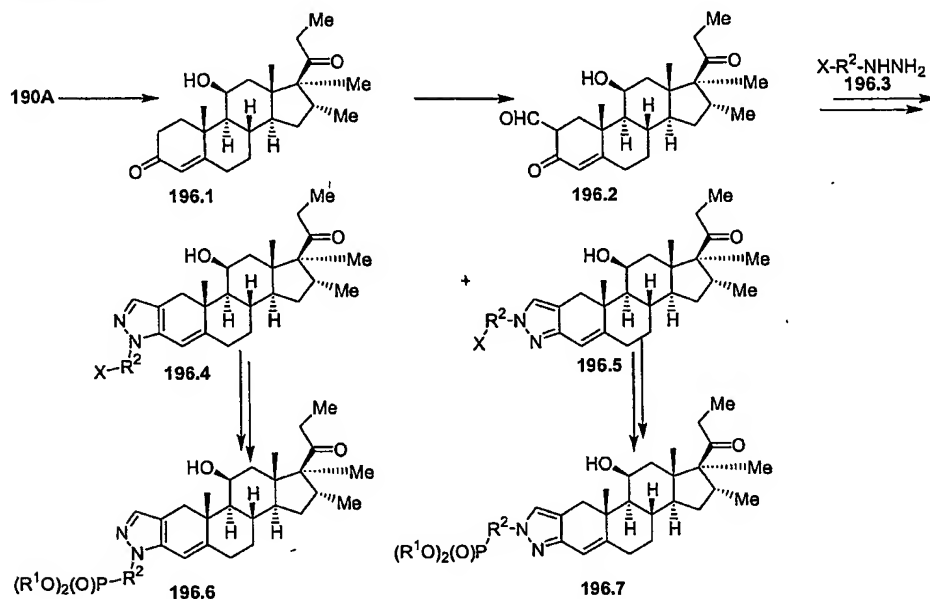
Alternatively, the carboxylic acid may first be converted into an activated derivative such as the acid chloride, anhydride, mixed anhydride, imidazolid and the like, and then reacted with the amine, in the presence of an organic base such as, for example, pyridine, to afford the amide.

The conversion of a carboxylic acid into the corresponding acid chloride can be effected by treatment of the carboxylic acid with a reagent such as, for example, thionyl chloride or oxalyl chloride in an inert organic solvent such as

dichloromethane, optionally in the presence of a catalytic amount of dimethylformamide.

Using the above procedures, but employing, in place of the carboxy-substituted hydroxylamine, **195.1**, different carboxy-substituted hydroxylamines, and/or different amino-substituted phosphonates, the products analogous to **195.4** are obtained.

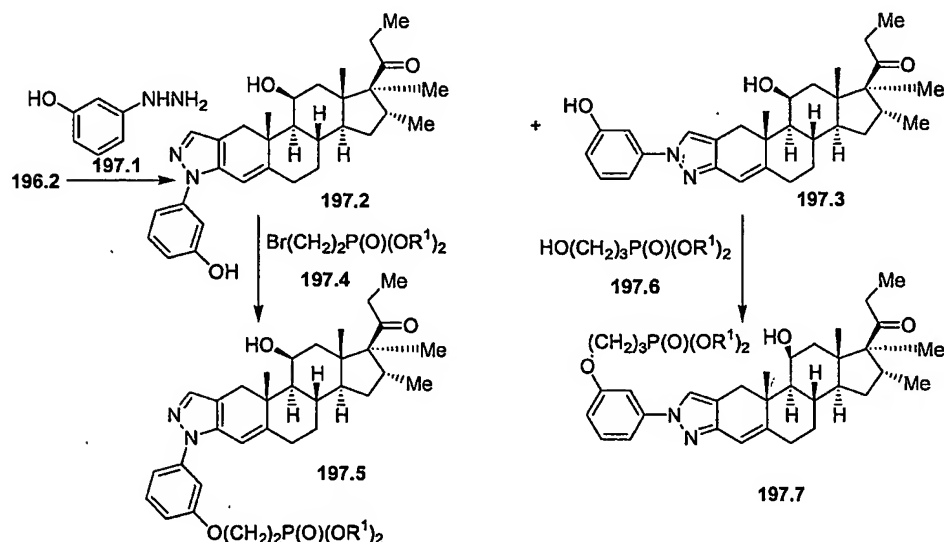
Example 196



The dienone, **190A**, is reduced to afford the 1,2-dihydro product, **196.1**. The catalytic hydrogenation reaction is effected by the use of tris(triphenylphosphine)rhodium (I) chloride, for example as described in *J. Med. Chem.*, **2001**, 44:, 602. The product is then reacted with ethyl formate and a base such as sodium hydride, in an inert solvent such as toluene or dimethylformamide, as described in *J. Am. Chem. Soc.*, **1964**, 86, 1520, to afford the 2-formyl product, **196.2**. This compound is then reacted with an alkyl, aralkyl, aryl or heteroaryl hydrazine, **196.3**, in which the substituent X is either a phosphonate group or a group which is subsequently transformed into a phosphonate-containing substituent. For example, X is dialkylphosphono, bromo, hydroxy, amino, carboxyl and the like. The reaction yields the isomeric 2'- and 1'-aryl pyrazoles, **196.4** and **196.5**. The pyrazole-forming reaction is performed between

equimolar amounts of the reactants in an acidic solvent such as acetic acid, as described in *J. Am. Chem. Soc.*, 1964, 86, 1520. The pyrazoles, 196.4 and 196.5, are then transformed, for example, by the procedures described in Examples 192 and 193, into the phosphonates, 196.6 and 196.7. Optionally, the reduction and formylation reactions are performed on the substrate, 190.1, in which the 20-ketone is protected as the cyclic ethylene ketal.

Example 197



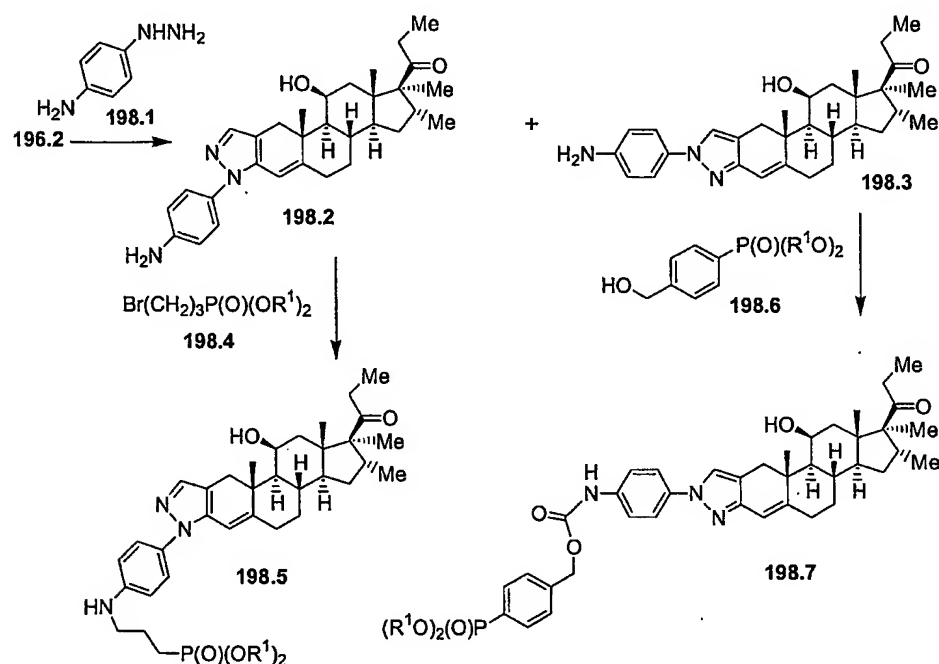
The ketoaldehyde, 196.2, is reacted, as described above, with 3-hydroxyphenyl hydrazine, 197.1, (JP 03011081) to give the pyrazoles, 197.2 and 197.3. The 2'-substituted isomer, 197.2, is then reacted in dimethylformamide solution at 70° with one molar equivalent of a dialkyl 2-bromoethyl phosphonate, 197.4, (Aldrich) and potassium carbonate, to give the ethoxy phosphonate, 197.5.

The isomeric pyrazole, 197.3, is reacted in a Mitsunobu with one molar equivalent of a dialkyl 3-hydroxypropyl phosphonate, 197.6, (*Zh. Obschei. Khim.*, 1974, 44, 1834) to yield the phosphonate, 197.7. The preparation of aromatic ethers by means of the Mitsunobu reaction is described, for example, in R. C. Larock, *Comprehensive Organic Transformations*, 448 (VCH, 1989), and in F.A. Carey and R. J. Sundberg, *Advanced Organic Chemistry*, Part B, 153-4 (Plenum, 2001) and in *Org. React.*, 1992, 42, 335. The phenol and the alcohol or thiol component are reacted together in an aprotic solvent such as, for

example, tetrahydrofuran, in the presence of a dialkyl azodicarboxylate and a triarylphosphine, to afford the ether or thioether products. The procedure is also described in *Org. React.*, 1992, 42, 335-656.

Using the above procedures, but employing different hydroxy-substituted hydrazines, and/or different bromo or hydroxy-substituted phosphonates, the products analogous to 195.5 and 197.6 are obtained.

Example 198



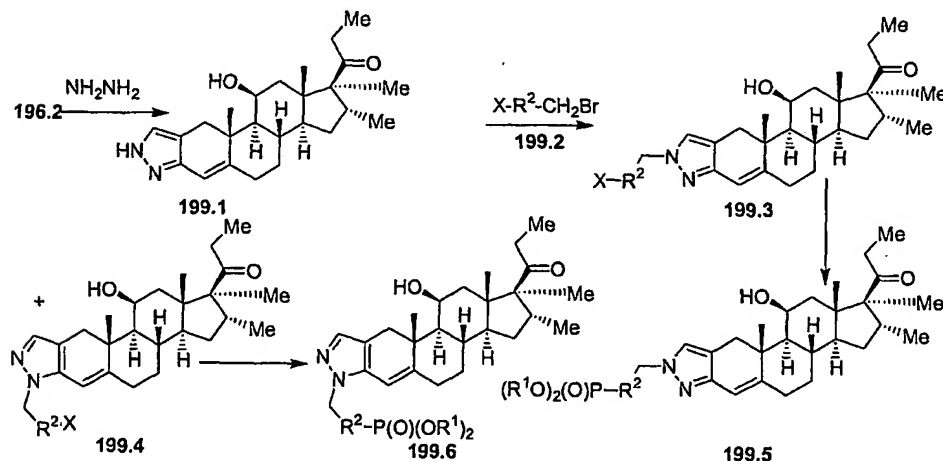
The ketoaldehyde, 196.2, is reacted, as described above, with 4-aminophenyl hydrazine, 198.1, (*Syn. Comm.*, 1974, 4, 57) to produce the pyrazoles, 198.2 and 198.3. The 2'-substituted isomer, 198.2, is then reacted in dimethylformamide solution at 70° with one molar equivalent of a dialkyl 3-bromopropyl phosphonate, 198.4, (*J. Amer. Chem. Soc.*, 2000, 122, 1554) and cesium carbonate, to give the amine phosphonate, 198.5.

Alternatively, the 1'-substituted pyrazole, 198.3, is coupled with a dialkyl 4-hydroxymethylphenyl phosphonate, 198.6, (U.S. Patent No. 5,569,664) and carbonyl diimidazole to prepare the carbamate phosphonate, 198.7. The preparation of carbamates is described in *Comprehensive Organic Functional*

Group Transformations, A. R. Katritzky, ed., Pergamon, 1995, Vol. 6, p 416ff, and in *Organic Functional Group Preparations*, by S. R. Sandler and W. Karo, Academic Press, 1986, p. 260ff. In the procedure, the amine is reacted in an inert aprotic solvent such as dichloromethane or tetrahydrofuran, with phosgene or a functional equivalent thereof, such as carbonyl diimidazole, triphosgene, pentafluorophenyl carbonate and the like, to afford the corresponding activated acylamine. The latter compound is then reacted with an alcohol to yield the carbamate.

Using the above procedures, but employing, in place of the aminophenyl hydrazine, 198.1, different amino-substituted hydrazines, and/or different dialkyl bromo or hydroxy-substituted phosphonates, the products analogous to the compounds, 198.5 and 198.7 are obtained.

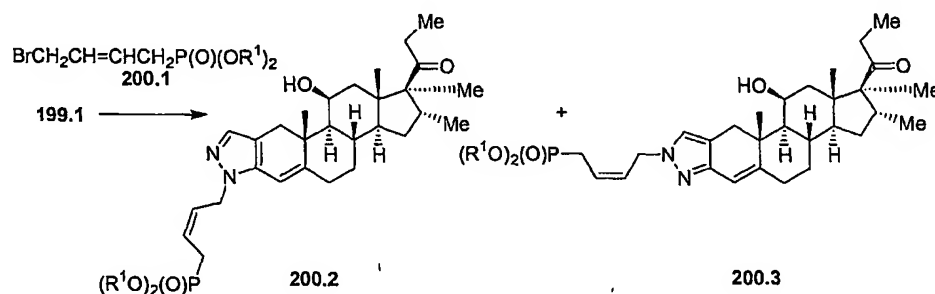
Example 199



The ketoaldehyde, 196.2, is reacted with hydrazine to afford the pyrazole derivative, 199.1. The reaction of steroidal 2-formyl-3-ketones with hydrazine is described in *J. Am. Chem. Soc.*, 1964, 86, 1520. The reaction is performed in acetic acid at ambient temperature. The pyrazole product is then reacted with a bromomethyl compound, 199.2, in which R^2 and X are as defined above, or a reactive bromoheteroaromatic reagent, to yield the alkylation products, 199.3 and 199.4. The alkylation of substituted pyrazoles is described, for example, in *Heterocyclic Chemistry*, by T. L. Gilchrist, Longman, 1992, p. 309. The reaction is performed between equimolar amounts of the substrates in a polar

solvent such as dimethylformamide or tetrahydrofuran, in the presence of a base such as dimethylaminopyridine, lithium hexamethyldisilazide and the like. The products, **199.3** and **199.4**, are, except in cases where *X* is dialkylphosphono, converted into the phosphonates, **199.5** and **199.6**, using the procedures described herein.

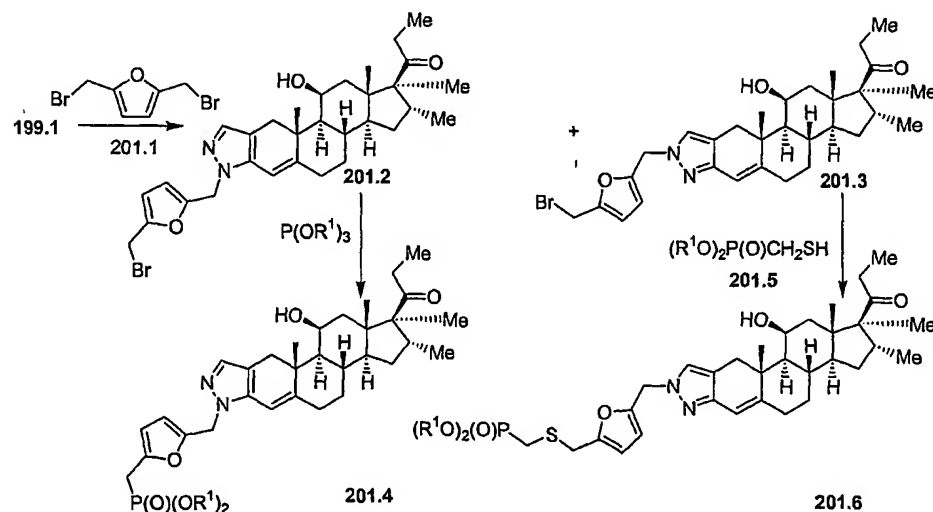
Example 200



The pyrazole, **199.1**, is reacted in dimethylformamide solution at 70°C with one molar equivalent of a dialkyl 4-bromobutenyl phosphonate, **200.1**, (*J. Med. Chem.*, **1992**, *35*, 1371) and lithium hexamethyl disilazide, to give the pyrazoles, **200.2** and **200.3**.

Using the above procedures, but employing different bromo-substituted phosphonates, the products analogous to **200.2** and **200.3** are obtained.

Example 201

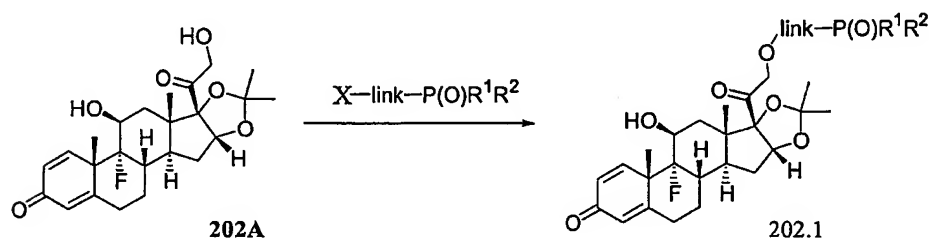


The pyrazole, 199.1, is reacted in tetrahydrofuran solution with 2,5-bis(bromomethyl)furan, 201.1, (*Tet.*, 1999, 55, 4709) and potassium hexamethyl disilazide, to give the alkylation products, 201.2 and 201.3. The 2'-substituted isomer, 201.2, is then reacted, in a Arbuzov reaction, with a trialkyl phosphite to yield the phosphonate, 201.4. The Arbuzov reaction is described in *Handb. Organophosphorus Chem.*, 1992, 115. In this procedure, in which a bromo substituent is converted into the corresponding phosphonate, the substrate is heated at from about 60° to about 160° with a five to fifty-fold molar excess of a trialkyl phosphite, to effect the transformation.

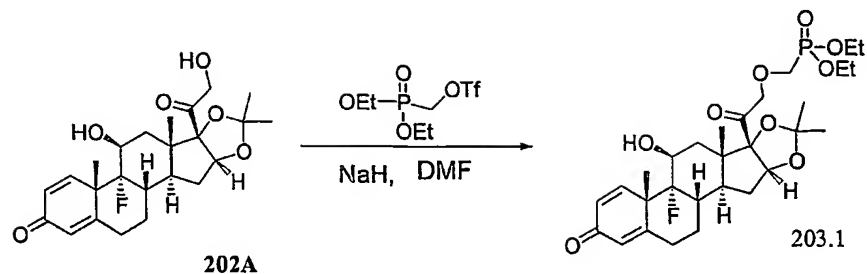
The 1'-substituted pyrazole, 201.3, is reacted at ambient temperature in dimethylformamide solution with one molar equivalent of a dialkyl mercapto-methyl phosphonate, 201.5, (*J. Med. Chem.*, 1985, 26, 1688) and cesium carbonate, to give the thioether phosphonate, 201.6.

Using the above procedures, but employing different dihalides, and/or different mercapto-substituted phosphonates, the products analogous to 201.4 and 201.6 are obtained.

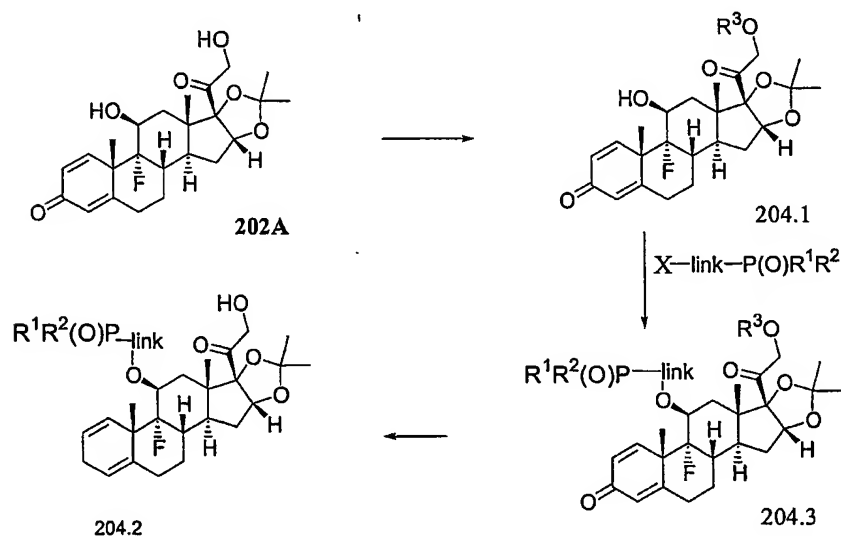
Example 202



Derivatives of the C-21 primary hydroxy group of the type, 202.1, are readily prepared by alkylating triamcinolone acetonide, 202A, with the appropriate phosphonate as shown.

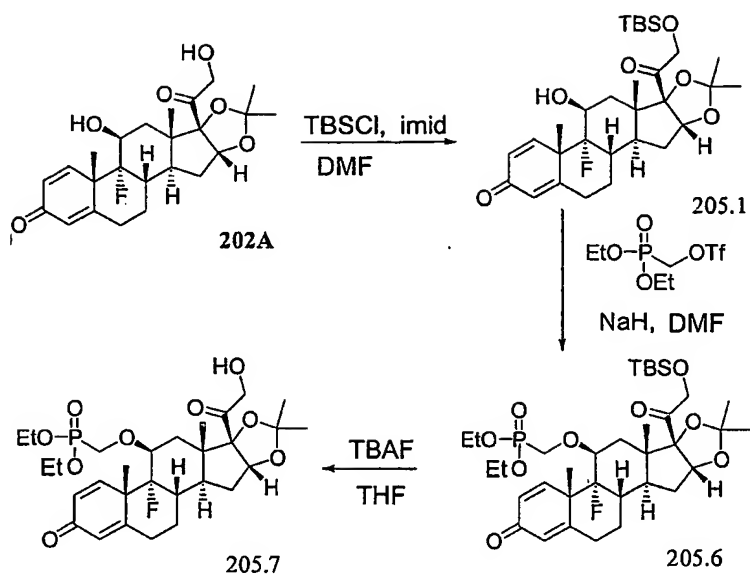
Example 203

After chemoselective extraction of the primary hydroxy proton of compound, **202A**, using one equivalent of sodium hydride, the phosphonate triflate is added to provide the ether, **203.1**.

Example 204

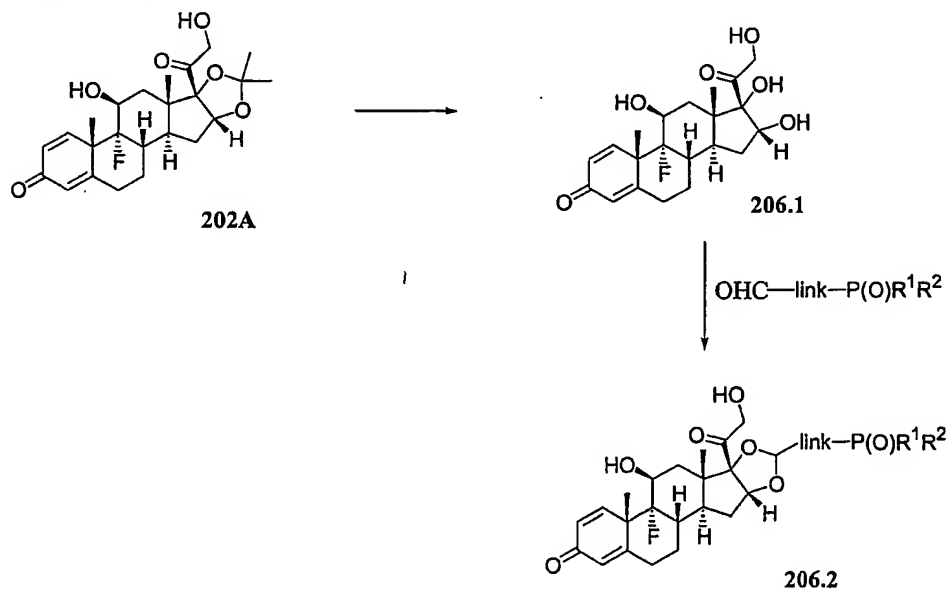
The primary hydroxy group is masked by an appropriate protecting group. After alkylation at the secondary hydroxy moiety of, **204.1**, with a leaving group-attached phosphonate and subsequent deprotection, desired analog, **204.2**, is obtained.

Example 205



Triamcinolone acetonide, **202A**, is chemoselectively protected as its silyl ether using the standard TBSCl and imidazole conditions (*J. Am. Chem. Soc.* **1972**, *94*, 6190). Alkylation at the exposed secondary hydroxy group with sodium hydride and the phosphonate triflate furnishes the intermediate, **205.6**. Final TBAF deprotection of the silyl ether affords the desired product, **205.7**.

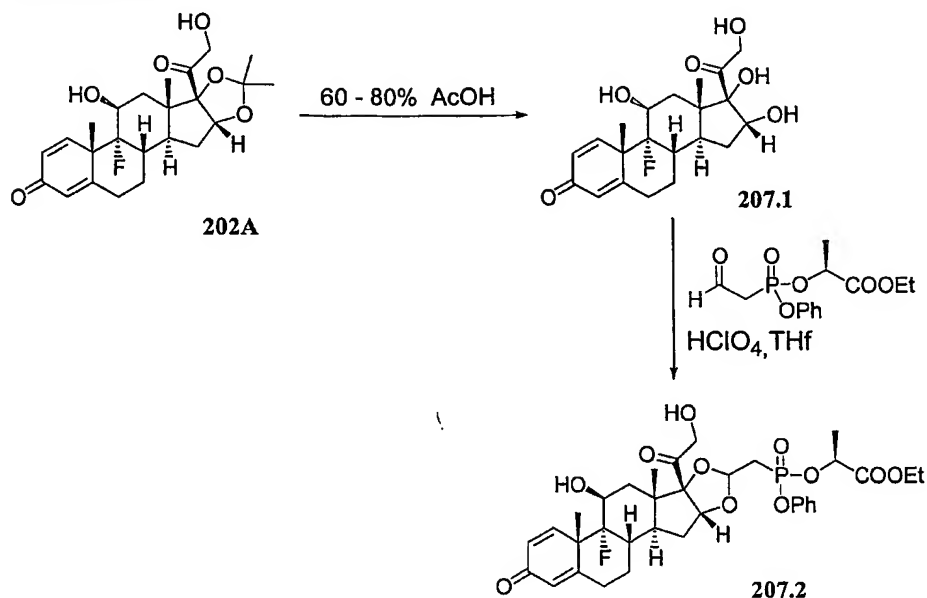
Example 206



Phosphonate derivatives of the acetal are readily prepared from acidic hydrolysis of triamcinolone acetonide, **202A**, to the diol, **206.1**, as illustrated in

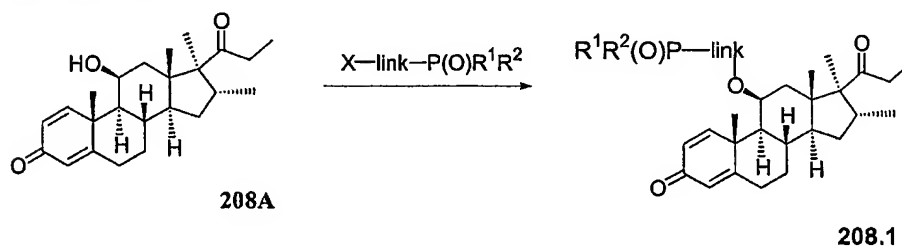
Scheme 3.1. Acetylation of the diol with a phosphonate aldehyde furnishes the desired acetal, **206.2**.

Example 207

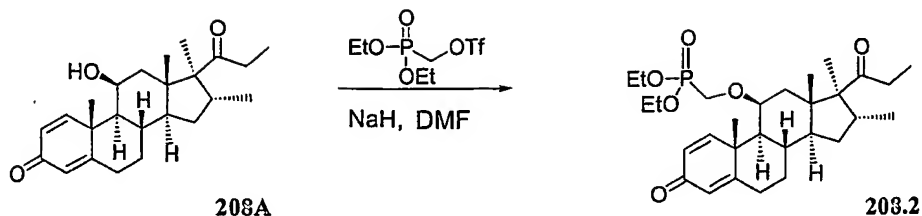


Triamcinolone acetonide, **202A**, is first hydrolyzed in aqueous acetic acid. (*Can. J. Chem.* 1983, 61, 634) The resulting diol, **207.1**, is acetalized with the phosphonate aldehyde and perchloric acid, affording the acetal, **207.2** (*J. Med. Chem.* 1996, 39, 4888-4896).

Example 208

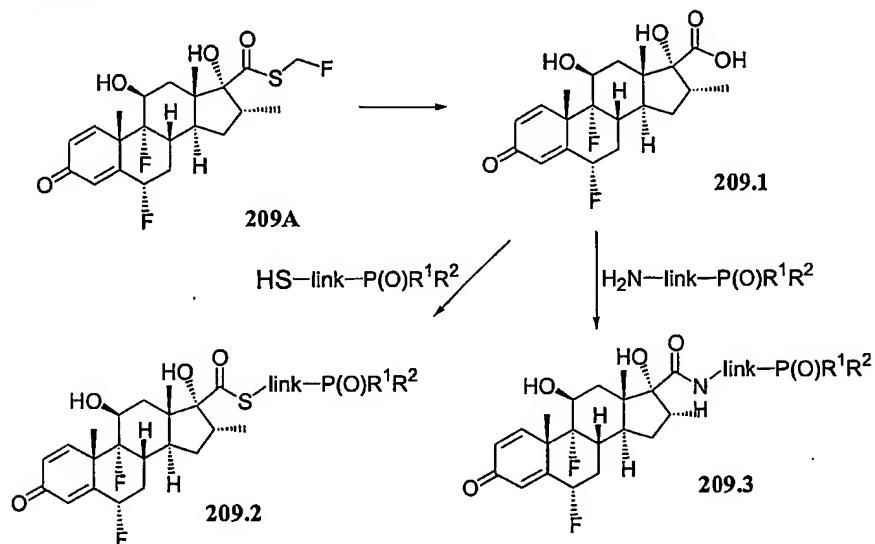


Derivatization at the C-11 hydroxy group is accomplished through alkylation of rimexolone, **208A**, with the appropriate phosphonate, furnishing analogs of the type, **208.1**.

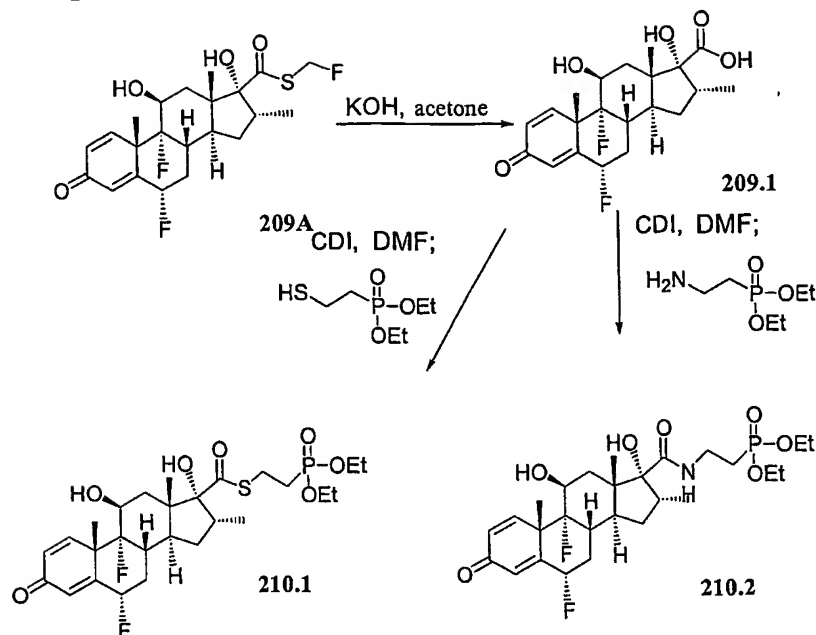


After sodium hydride extraction of the hydroxy proton in, **208A**, diethyl phosphonate triflate is added to afford ether, **208.2**.

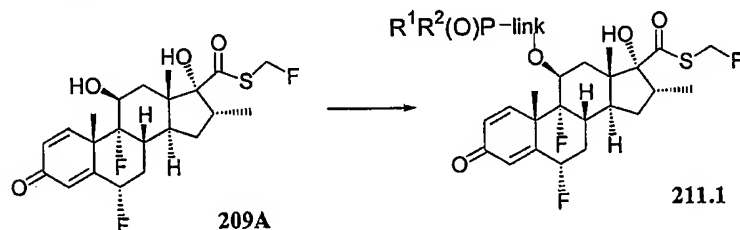
Example 209



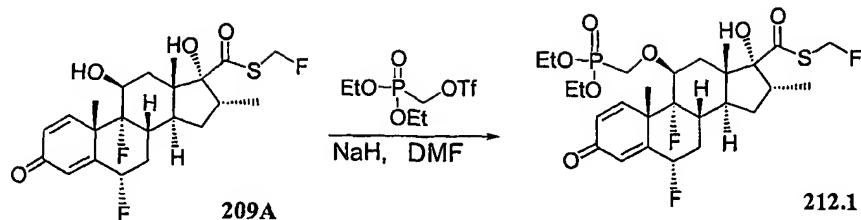
Derivatives of the carbonyl at C-17 are readily prepared from saponification of fluticasone, **209A**, to the carboxylic acid, **209.1**. Activation of the carboxylic acid, followed by reaction with thiophosphonate or aminophosphonate nucleophile furnishes the desired thioester, **209.2**, and amide, **209.3**, respectively.

Example 210

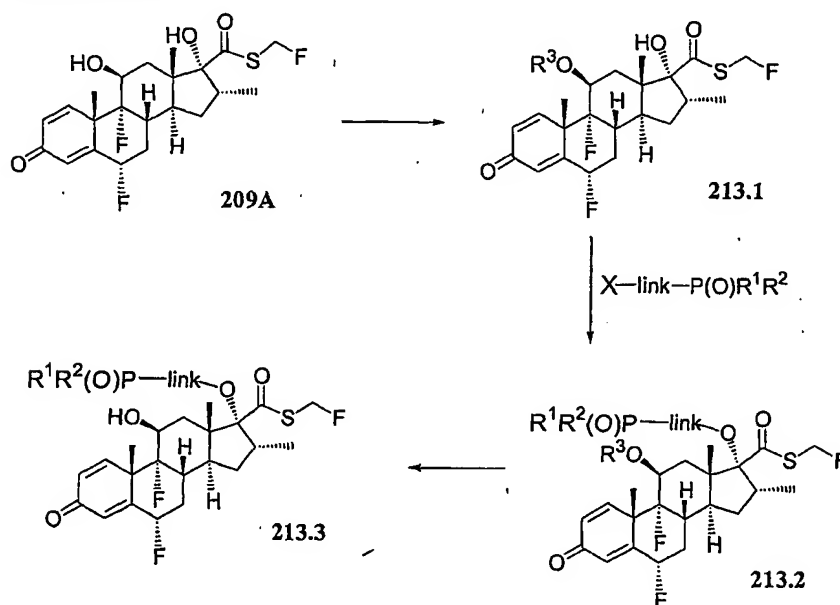
Fluticasone, **209A**, is first saponified with potassium hydroxide in acetone. (*Synthesis*, **2002**, 921-927) The resulting carboxylic acid, **209.1**, is activated to the carboxylic acid imidazole by the addition of 1,1'-carbonyldiimidazole (CDI) (*J. Med. Chem.* **1994**, 37, 3717-3729). Treatment with the thiophosphonate affords thioester, **210.1**. Magnesium ethoxide may be added to help enhance the reactivity (*Tetrahedron Lett.* **1981**, 22, 3245-3246). Alternatively, the carboimidazole intermediate derived from, **209.1**, can be reacted with the aminophosphonate to produce amide, **210.2**.

Example 211

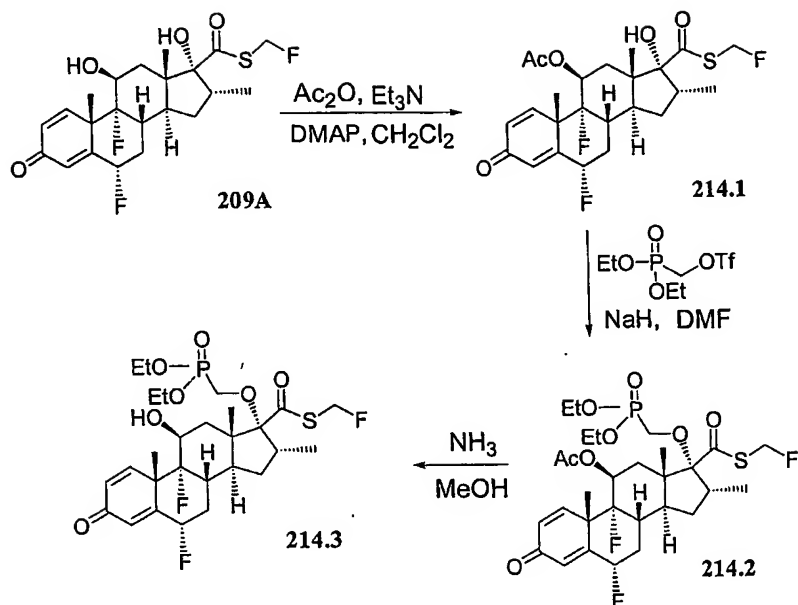
The less sterically hindered C-11 hydroxy group is selectively alkylated with the appropriate phosphonate to give analogs of formula, **211.1**.

Example 212

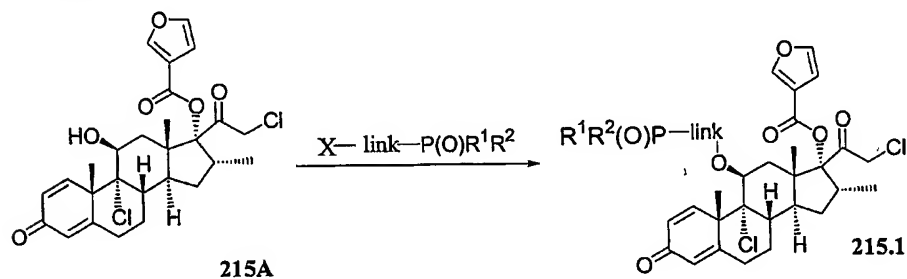
After regioselective extraction of the C-11 hydroxy proton in, **209A**, using one equivalent of sodium hydride, the phosphonate triflate is added to provide the ether, **212.1**.

Example 213

The C-11 hydroxy group is masked by an appropriate protecting group. After alkylation at the C-17 hydroxy moiety of, **213.1**, with a leaving group-attached phosphonate and subsequent deprotection, desired analog, **213.3**, is obtained.

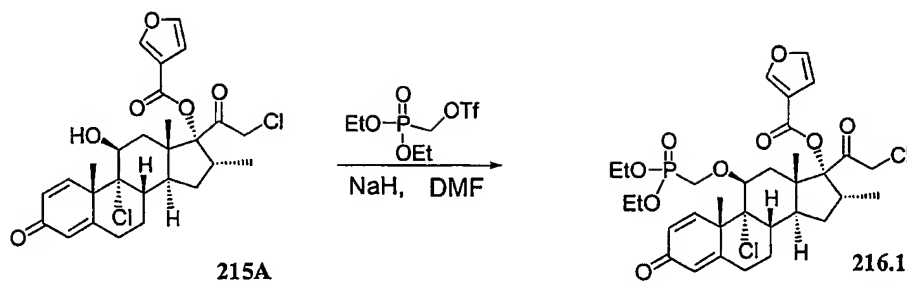
Example 214

Fluticasone **209A** is regioselectively protected as its C-11 acetate ester using the standard acetic anhydride and DMAP conditions (*J. Org. Chem.* 1998, 63, 2342-2347). Alkylation at the exposed C-17 hydroxy group with sodium hydride and the phosphonate triflate furnishes the intermediate, **214.2**. Final ammonia deprotection of the acetate affords the desired ether, **214.3**.

Example 215

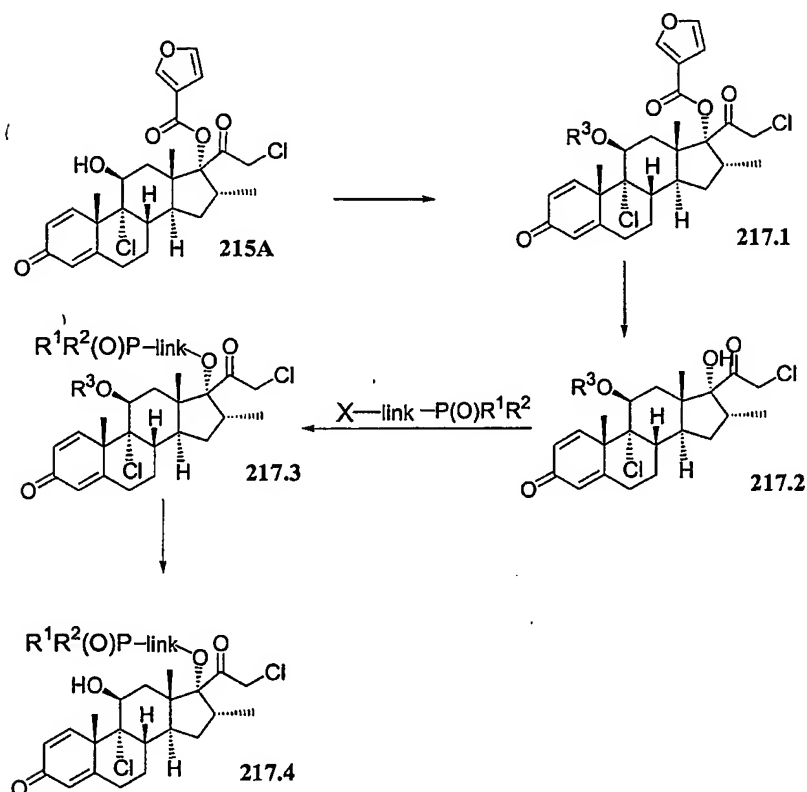
Derivatization at the C-11 hydroxy group is accomplished through alkylation of mometasone fuorate, **215A**, with the appropriate phosphonate, furnishing analogs of the type, **215.1**.

Example 216

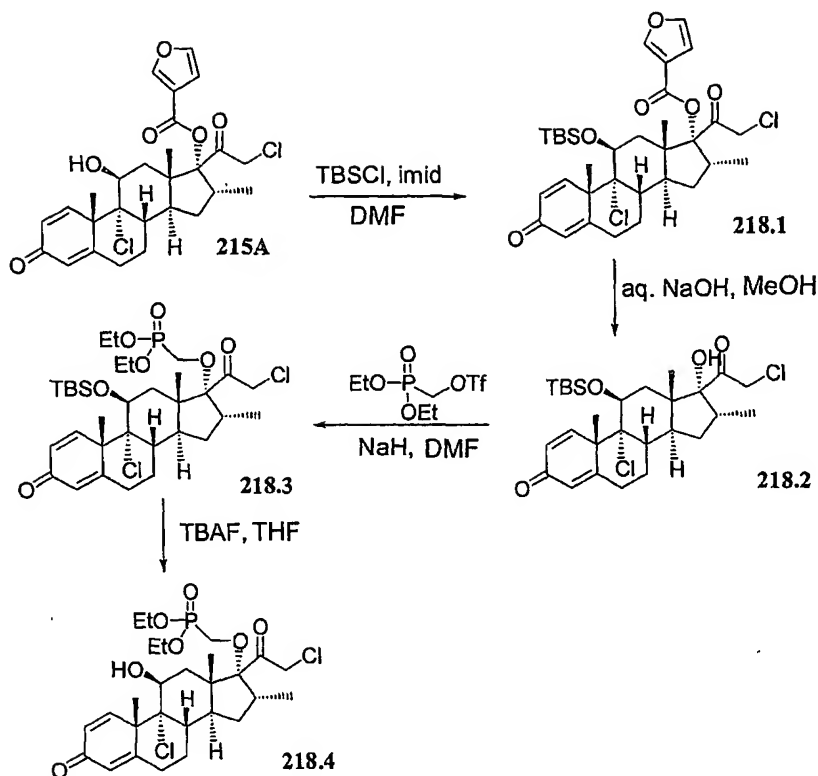


After sodium hydride extraction of the hydroxy proton in **215A**, diethyl phosphonate triflate is added to afford ether, **216.1**.

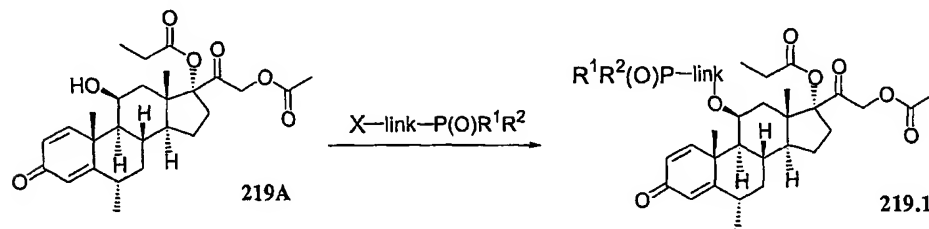
Example 217



Following protection of the only exposed hydroxy group in mometasone fuorate, **215A**, intermediate, **217.1**, is saponified to give alcohol, **217.3**. Alkylation at the C-17 hydroxy group with the appropriate phosphonate and subsequent deprotection provides the desired product, **217.4**.

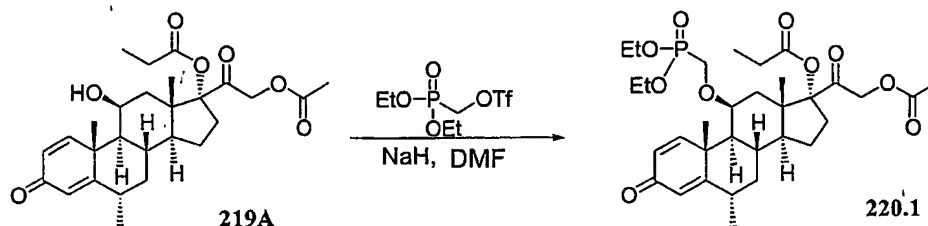
Example 218

Mometasone fuorate, **215A**, is protected as its silyl ether using the standard TBSCl and imidazole conditions (*J. Am. Chem. Soc.*, **1972**, *94*, 6190). Saponification of the fuoyl ester moiety using aqueous sodium hydroxide provides the alcohol, **218.1** (*J. Chem. Soc. Perkin Trans. 1*, **1993**, *12*, 1359-1366). The tertiary hydroxy group is alkylated by the addition of sodium hydroxide and the phosphonate triflate. After deprotection of the silyl ether in intermediate, **218.3**, with TBAF, diethyl phosponate, **218.4**, results.

Example 219

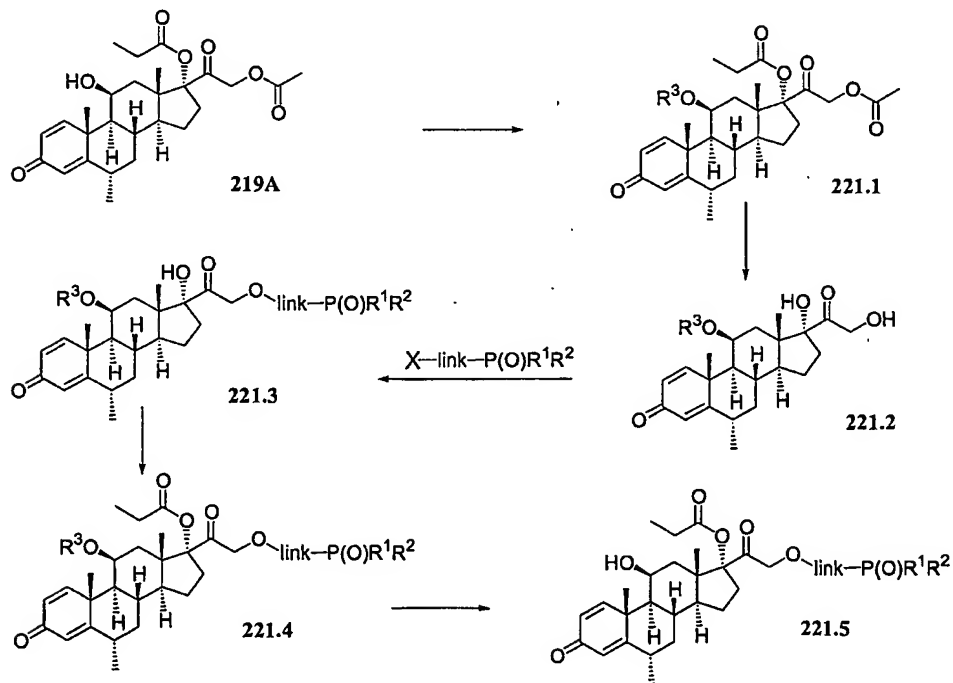
Derivatization at the C-11 hydroxy group is accomplished through alkylation of methylprednisolone aceponate (**219A**) with the appropriate phosphonate, furnishing analogs of Example 219.1.

Example 220



After sodium hydride extraction of the hydroxy proton in **219A**, diethyl phosphonate triflate is added to afford ether, **220.1**.

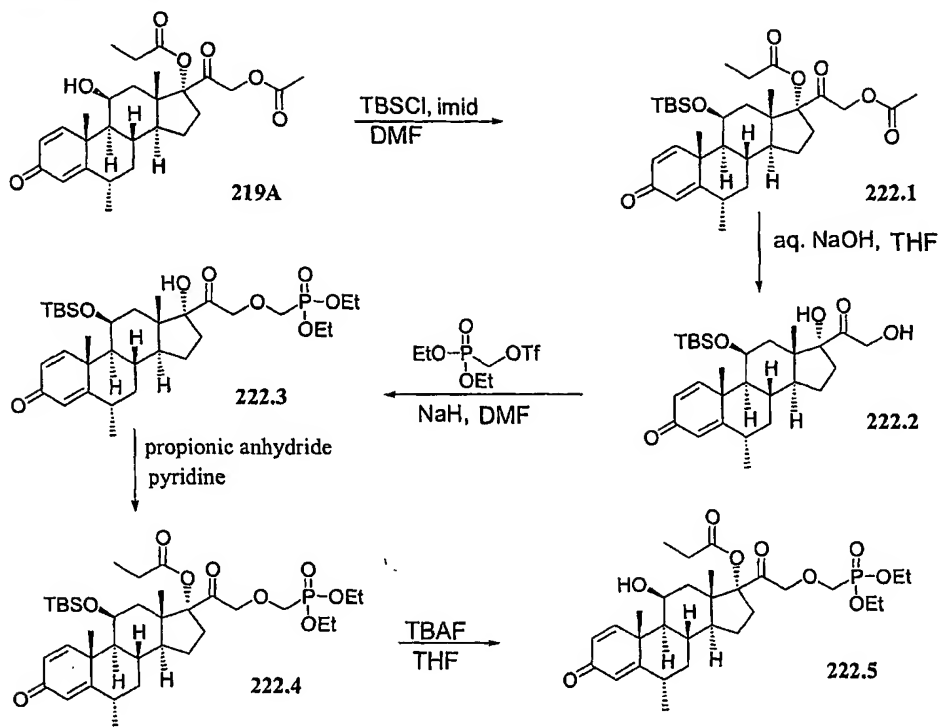
Example 221



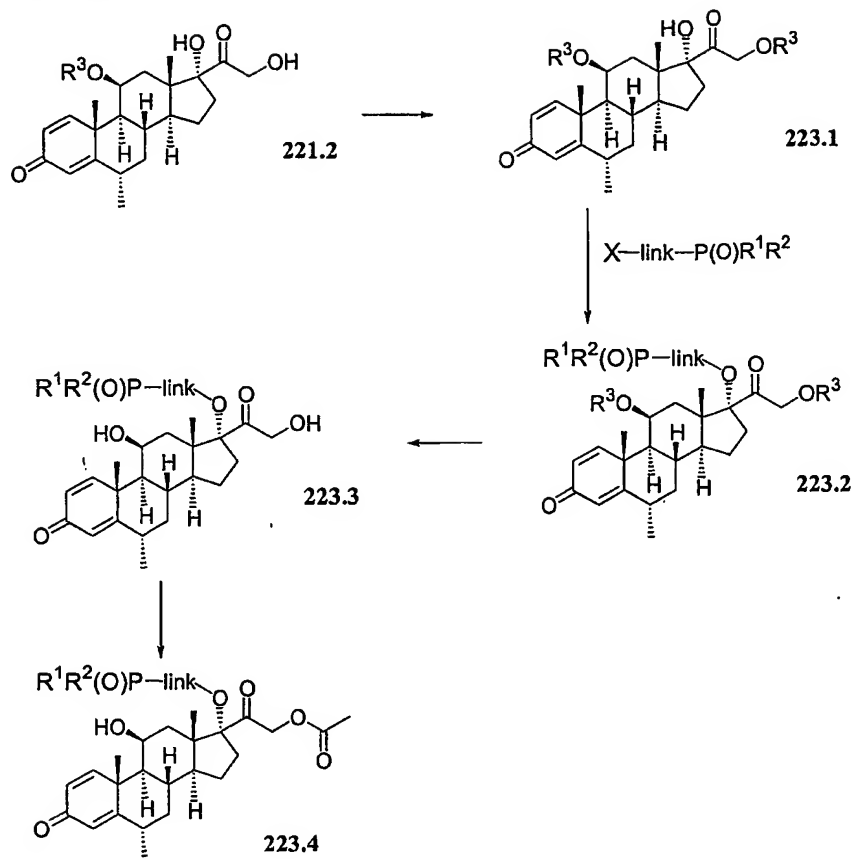
Following protection of the only exposed hydroxy group in **219A**, intermediate, **221.1**, is saponified to give diol, **221.2**. Alkylation at the primary hydroxy group with the appropriate phosphonate and subsequent acylation

provides the propionate ester, **221.4**. The desired product, **221.5**, is achieved after deprotection.

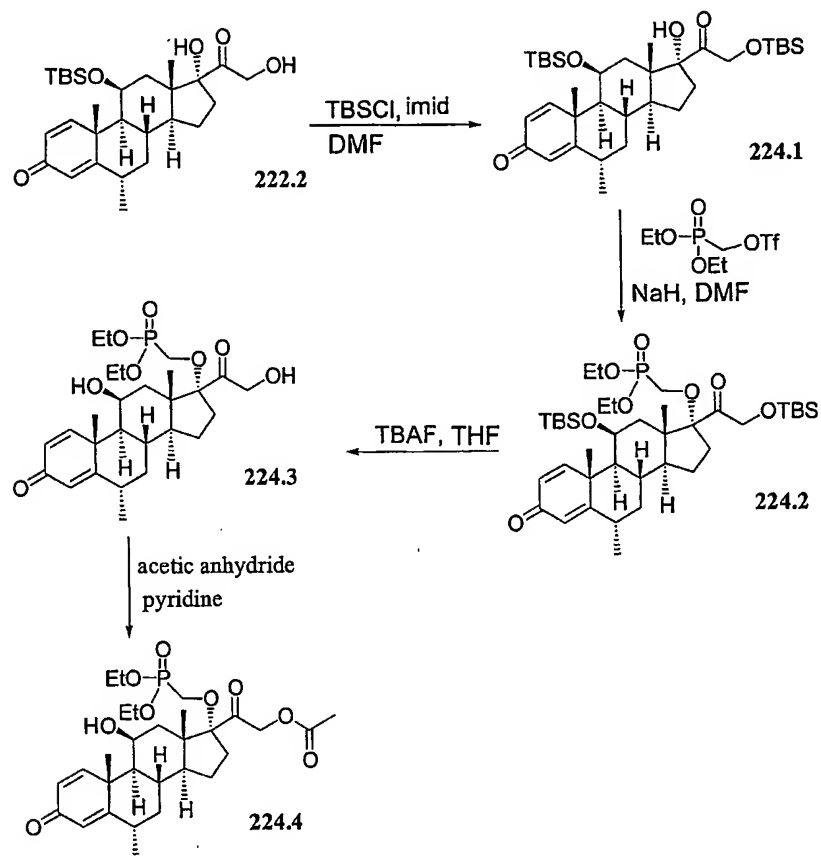
Example 222



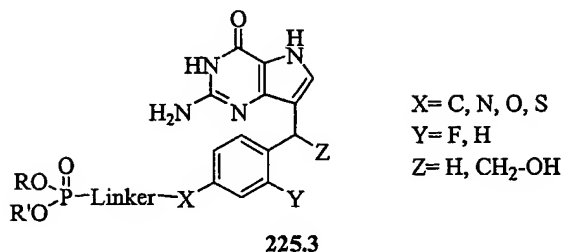
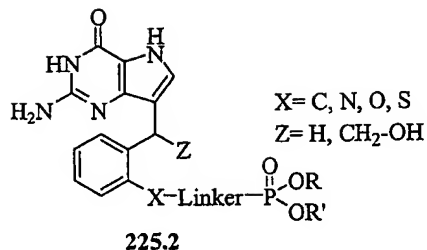
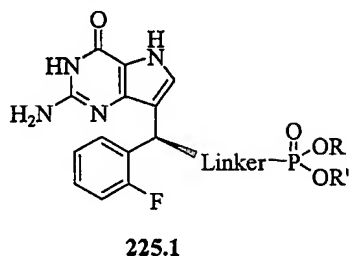
Methylprednisolone aceponate, **219A**, is protected as its silyl ether using the standard TBSCl and imidazole conditions (*J. Am. Chem. Soc.*, 1972, 94, 6190). Saponification of both ester moieties using aqueous sodium hydroxide provides the diol, **222.2**. The less sterically hindered primary hydroxy group is alkylated by the addition of sodium hydroxide and the phosphonate triflate. After treating intermediate, **222.3**, with propionic anhydride in pyridine, the previously hydrolyzed C-17 propionic ester is replaced (*J. Med. Chem.*, 1980, 23, 430-437). TBAF deprotection of the silyl ether furnishes diethyl phosphonate, **222.5**.

Example 223

The two hydroxy groups of diol, **221.2**, are regioselectively differentiated by protection at the primary site, thus allowing alkylation at the tertiary hydroxy group. The resulting phosphonate intermediate, **223.1**, is then deprotected to afford the diol, **223.3**. Again the more accessible primary hydroxy group is acylated to produce the desired analog, **223.4**.

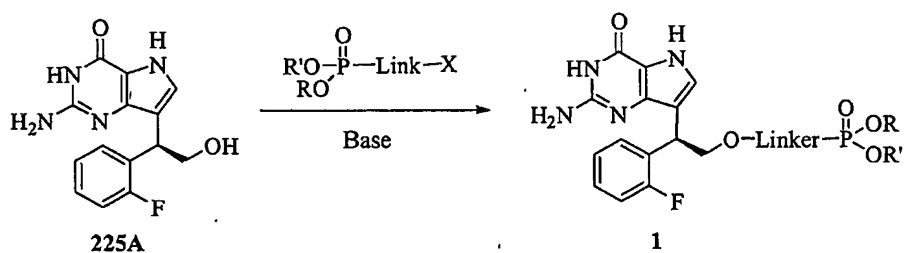
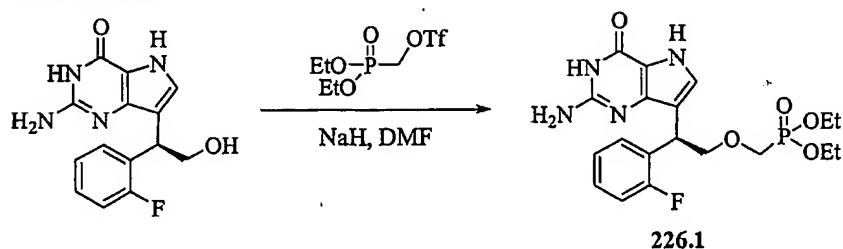
Example 224

Diol, 222.2, is alkylated with the diethyl phosphonate triflate, the resulting intermediate, 224.2, is treated with TBAF to give diol, 224.3. Acetic anhydride and pyridine are used to generate the final product, 224.4 (*J. Mol. Biol.*, 1972, 72, 219).

Example 225

In all cases, linker is 0-8 and preferably 1-6 atoms

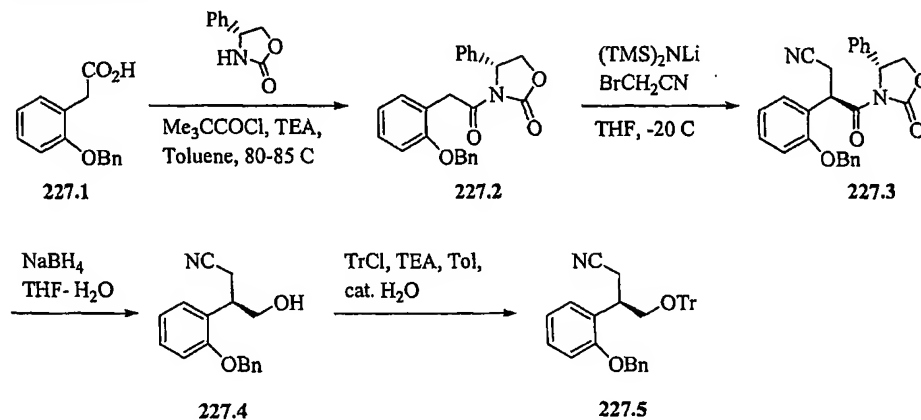
Compounds such as, **225.1**, can be made according to the general route outlined below.

**Example 226**

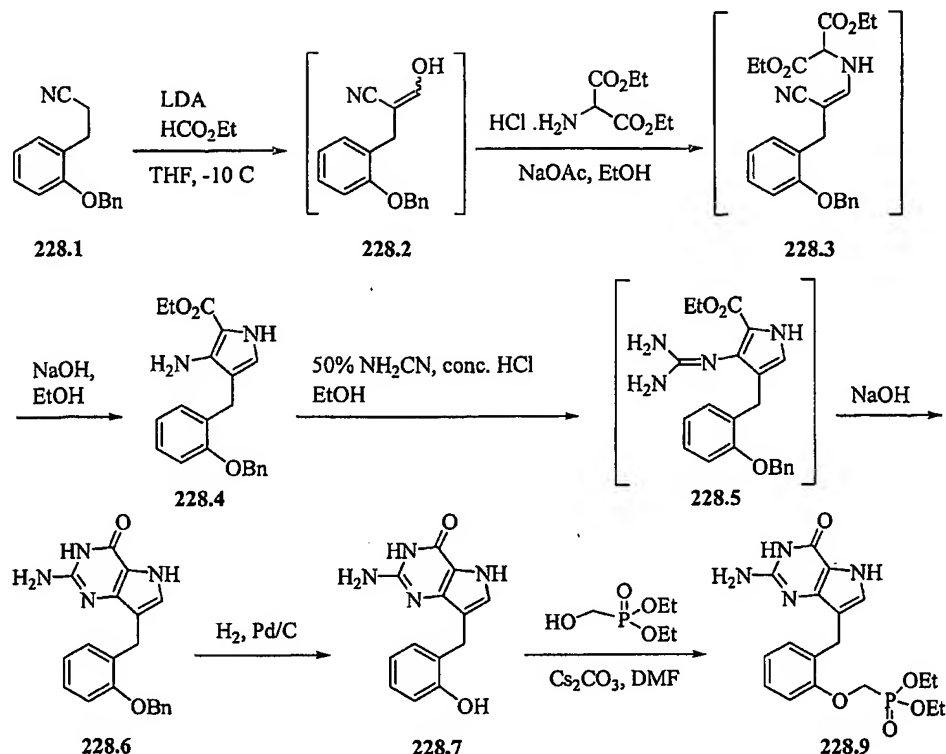
PNP-405, **225A**, is prepared according to the method of Littler, B. J. et al., 7th International Conference on Organic Process Research and Development, New Orleans, LA, March 16-19, 2003. PNP-405 is treated in a solvent such as tetrahydrofuran or dimethylformamide with a base such as

sodium hydride. When bubbling ceases, diethyl phosphonomethyltriflate (prepared according to *Tetrahedron Lett.*, 1986, 27, 1477) is added, to provide compound, **226.1**, as the desired product.

Example 227



Compounds such as **227.1** (where X= O, Z= CH_2OH), can be prepared according to the procedure of Littler, B. J. et al., 7th International Conference on Organic Process Research and Development, New Orleans, LA, March 16-19, 2003 (Schemes 3 and 4). The starting material, 2-benzyloxyphenylacetic acid, **227.1**, (provided by Avocado) can be acylated via the mixed anhydride with the oxazolidinone shown at 80-85 °C, with triethylamine as base to provide compound, **227.2**. A low-temperature alkylation with bromoacetonitrile results in the formation of compound, **227.3**, with good diastereomeric ratio. Removal of the chiral auxiliary under reductive conditions yields compound, **227.4**, without racemization. Protection of the resulting alcohol with the trityl group provides compound, **227.5**. Subsequent pyrrole ring construction as well as cyclo-guanidinylation reaction to prepare the six-membered 2-aminopyrimidone ring is performed as described in Example 228 for compound, **228.9**, to provide compound, **227.6**.

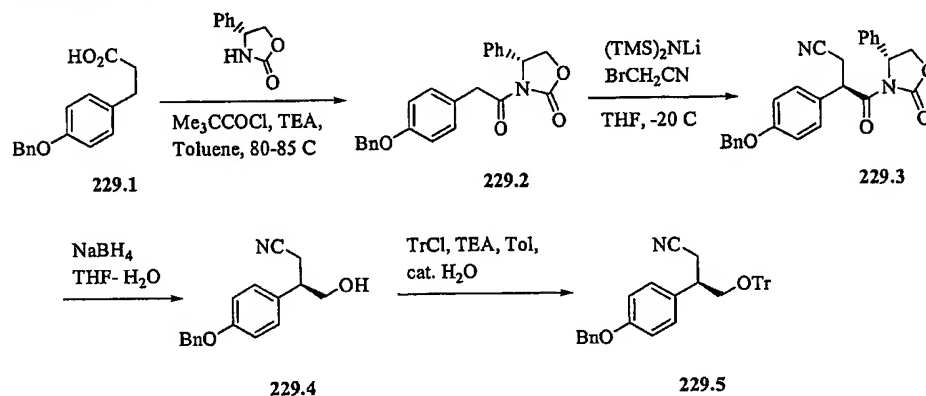
Example 228

Compound, **228.1**, (where $\text{X} = \text{O}$, $\text{Z} = \text{H}$) can be prepared according to the general route outlined above. The starting material, 3-(2-benzyloxy-phenyl)-propionitrile, **228.1**, is available by Lewis acid-mediated reaction of phenol with acrylonitrile according to U.S. Patent No. 2,789,995, published in 1954. Intermediate, **227.5**, can follow the same synthetic steps as outlined here to provide compound, **227.1**.

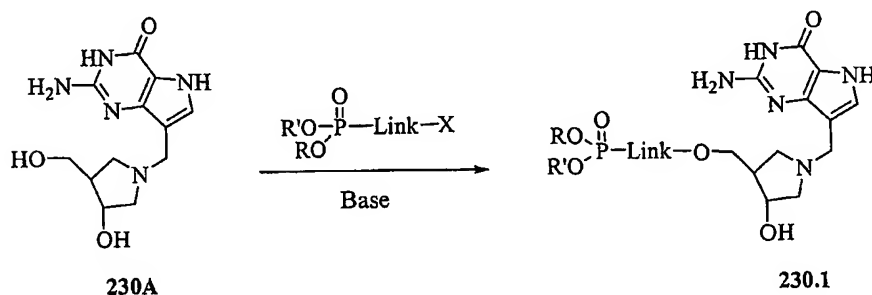
Pyrrole ring construction can be completed in three steps from 3-(2-benzyloxy-phenyl)-propionitrile, **228.1**. Formation of 3-hydroxy-acrylonitrile, **228.2**, can be achieved by exposure of, **228.1**, to LDA and ethyl formate. Condensation of this product with 2-Amino-malonic acid diethyl ester in EtOH and sodium acetate yields compound, **228.3**, which undergoes a decarboxylative cyclization in the basic medium of NaOH and EtOH to provide pyrrole, **228.4**. In case of compound, **227.1**, synthesis, the trityl protecting group on the benzylic alcohol is removed at this stage. Subsequently, guanidinylation reaction using cyanamide provides compound, **228.5**, which, upon treatment with sodium hydroxide, cyclizes to form the 2-aminopyrimidone ring (compound, **228.6**). Removal of the phenolic protecting group under hydrogenolysis conditions

provides the free phenol, which is used as the attachment site for the pro-drug group. A variety of linkers may be utilized to attach the pro-drug moiety to the backbone molecule. A particular example in which diethyl phosphonomethyltriflate is used as the starting materials is shown is above. Therefore, compound, **228.7**, is treated in a solvent such as tetrahydrofuran or dimethylformamide with a base such as sodium hydride or cesium carbonate. When bubbling ceases, diethyl phosphonomethyltriflate (prepared according to *Tetrahedron Lett.*, 1986, 27, 1477) is added, to provide compound, **228.2**, as the desired product.

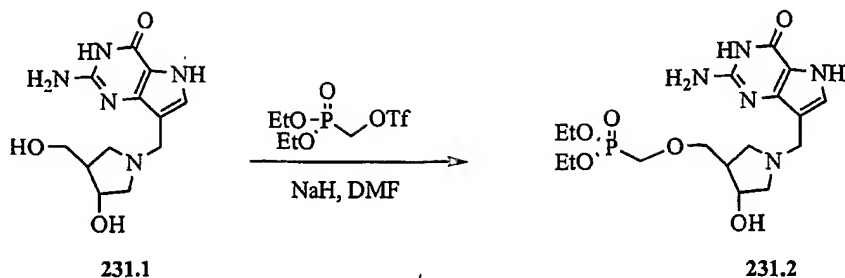
Example 229



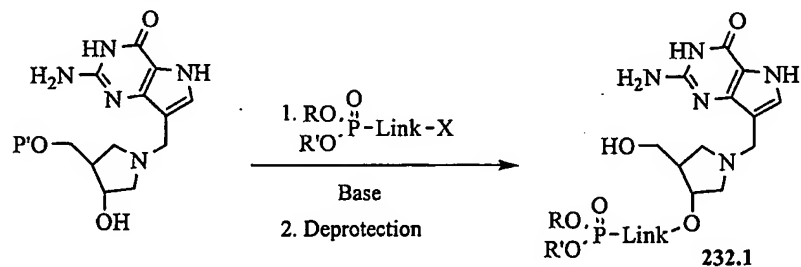
Compounds such as, **229.1**, (where X= O, Y= H, Z= CH_2OH) can be prepared from 4-benzyloxyphenylacetic acid (available from Aldrich). Following a similar sequence to that demonstrated above, intermediate, **229.5**, can be prepared. Proceeding with the sequence shown above, **229.5**, can be transformed to the desired product, **229.1**.



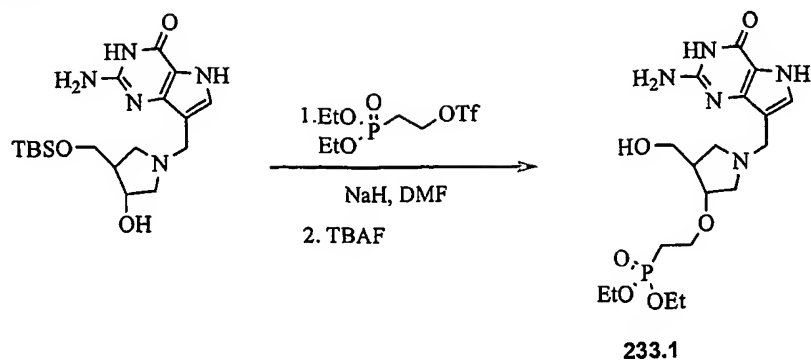
Compounds such as, **230A**, can be made according to the general route outlined above.

Example 231

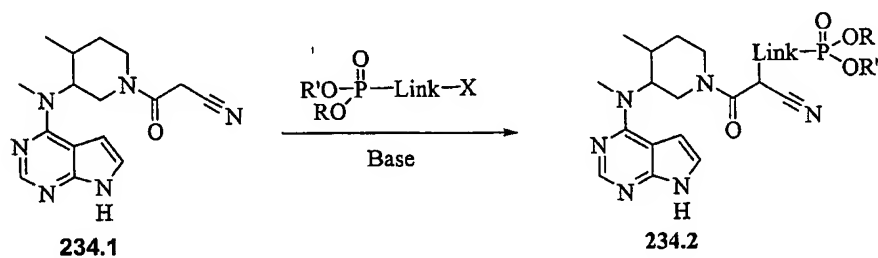
Preparation of DADMe-ImmG is reported in Lewandowics A. et al., *Biochemistry*, **2003**, *42*, 6057. The tertiary nitrogen of the ring may not interfere with the alkylation of the secondary alcohol and in that case does not need to be protected, although standard protection and deprotection protocols as described in Greene, T., *Protective Groups in Organic Synthesis*, Wiley-Interscience, **1999** may be used if necessary. Reaction of the primary alcohol, **231.1**, with base followed by addition of the appropriately activated phosphonate yields the protected product. Global deprotection yields the desired phosphonate, **231.2**.

Example 232

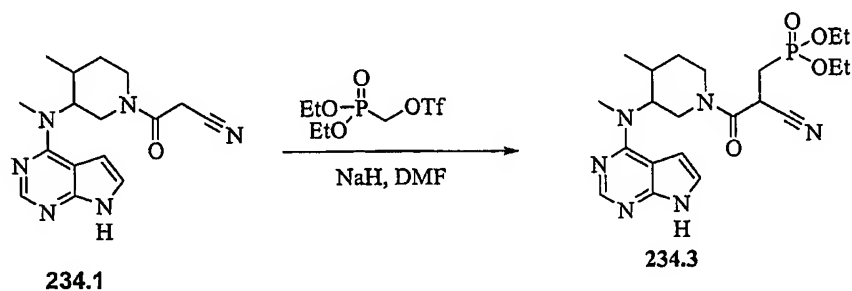
Compounds such as, **232.1**, can be made according to the general route outlined above.

Example 233

The protected DADMe derivative can be treated with treated in a solvent such as tetrahydrofuran or dimethylformamide with a base such as sodium hydride. When bubbling ceases, diethyl phosphonoethyltriflate (prepared according to *Tetrahedron Lett.*, 1986, 27, 1477) is added, yielding the desired phosphonate ester. Removal of the protecting group can be performed as described in Greene, T., *Protective Groups in Organic Synthesis*, Wiley-Interscience, 1999 to provide the desired phosphonate ester, **233.1**.

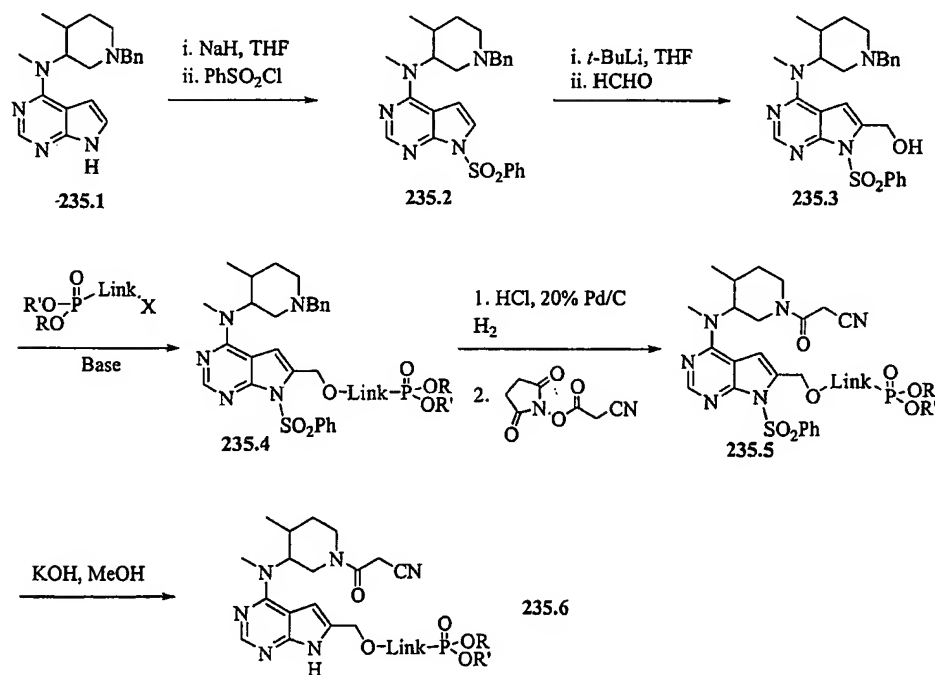
Example 234

Compounds such as, **234.2**, can be made according to the general route outlined above.



CP-690,550, 3-{4-methyl-3-[methyl-(7H-pyrrolo[2,3-d]pyrimidin-4-yl)-amino]-piperidin-1-yl}-3-oxo-propionitrile, **234.1**, can be prepared as described in WO 02096909 and WO 03048162. Enolate formation at the α -cyanoamide position using over 2 equivalents of base followed by addition of diethyl phosphonomethyltriflate (prepared according to *Tetrahedron Lett.*, 1986, 27, 1477) yields the desired compound, **234.3**. A solvent such as THF, DMF or other anhydrous solvents may be used for this reaction. In case the pyrrole nitrogen interferes with the desired alkylation, a protecting group such as BOC may be introduced before the alkylation reaction. Removal of the BOC group can be accomplished by exposure of the reaction product to TFA as described in Greene, T., *Protective groups in Organic Synthesis*, Wiley-Interscience, 1999.

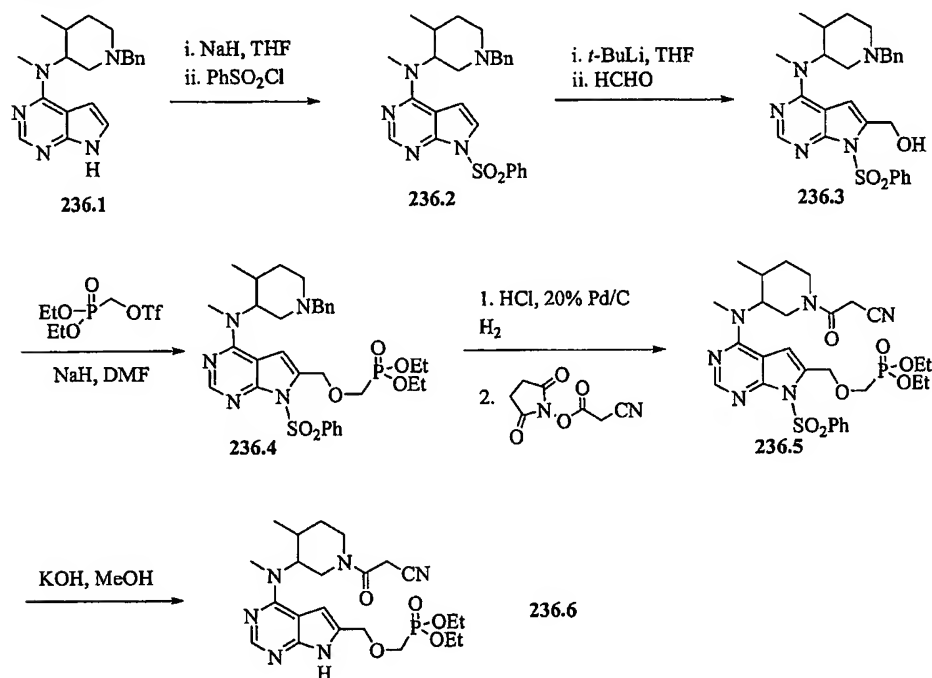
Example 235



Compound, **235.1**, is prepared according to WO 02096909. Protection of the pyrrole nitrogen using a tosyl group is achieved as described in Sakamoto, T. *et al.*, *Tetrahedron Lett.* 1994, 35, 18, 2919. Ortho lithiation using *t*-BuLi and quenching with formaldehyde as described in the above reference as well as Seela, F. *et al.*, *Chem. Ber.* 1977, 110, 4, 1462 introduces a substituent at the

requisite site. The primary alcohol so formed may be used for attachment of the phosphonate moiety via ether formation using base and diethyl phosphonomethyltriflate (prepared according to *Tetrahedron Lett.*, 1986, 27, 1477) in an anhydrous solvent. Removal of the benzyl protecting group is achieved using hydrogenolysis conditions. The piperidine nitrogen is then coupled with cyanoacetic acid 2,5-dioxo-pyrrolidine-1-yl ester to provide compound, 235.5. Removal of the tosyl protecting group can be achieved using basic conditions to provide the desired product, 235.6.

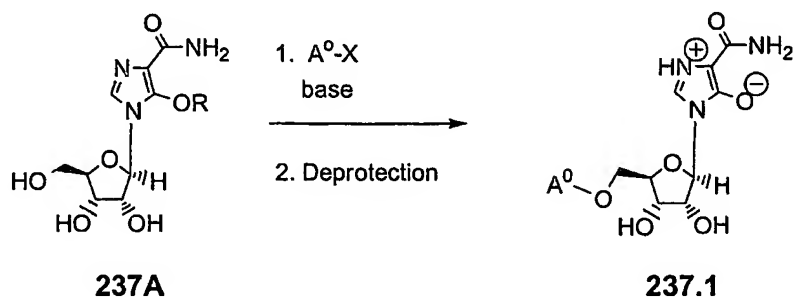
Example 236



Specifically, (1-benzyl-4-methyl-piperidin-3-yl)-methyl-(7H-pyrrolo[2,3-d]pyrimidin-4-yl)-amine, compound, 236.1, (prepared as described in WO 02,096,909) is first protected on the pyrrole nitrogen using a tosyl group. Subsequent formylation using the procedure reported by Sakamoto, T. *et al.*, (*Tetrahedron Lett.* 1994, 35, 2919) provides compound, 236.3. The primary alcohol is then treated in a solvent such as tetrahydrofuran or dimethylformamide with a base such as sodium hydride. When bubbling ceases, diethyl phosphonomethyltriflate (prepared according to *Tetrahedron Lett.*, 1986, 27, 1477) is added, yielding the desired product, 236.4. Debenzylation of the

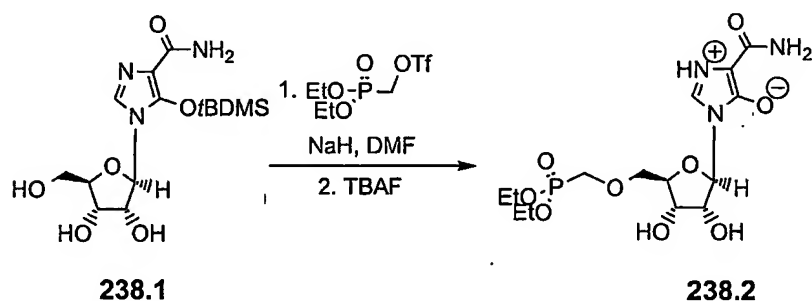
piperidine nitrogen following by coupling to cyano-acetic acid 2,5-dioxo-pyrrolidine-1-yl ester gives compound, **236.5**. Removal of the tosyl protecting group provides the desired pro-drug, **236.6**.

Example 237



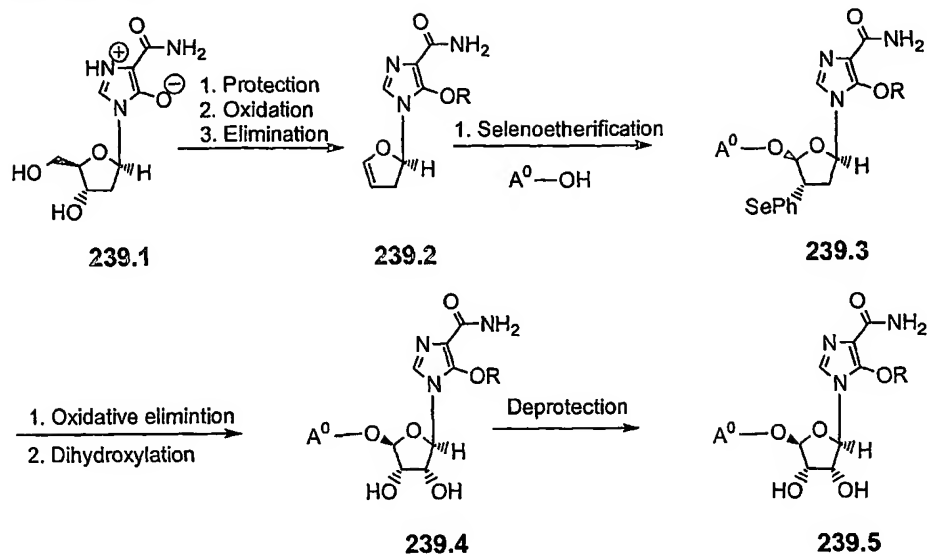
The syntheses of phosphonate compounds of the invention and of intermediate compounds necessary for their synthesis are illustrated herein.

Example 238



The preparation of phosphonates of phosphonate compounds of the invention is illustrated above. The 5-hydroxy-1- β -D-ribofuranosyl-1H-imidazole-4-carboxamide, **238.1** (prepared according to U.S. Patent No. 3,888,843), can be treated in a solvent such as tetrahydrofuran or dimethylformamide with a base such as sodium hydride. When bubbling ceases, diethyl phosphonomethyltriflate (prepared according to *Tetrahedron Lett.*, 1986, 27, 1477) is added, yielding the desired phosphonate diester **238.2**.

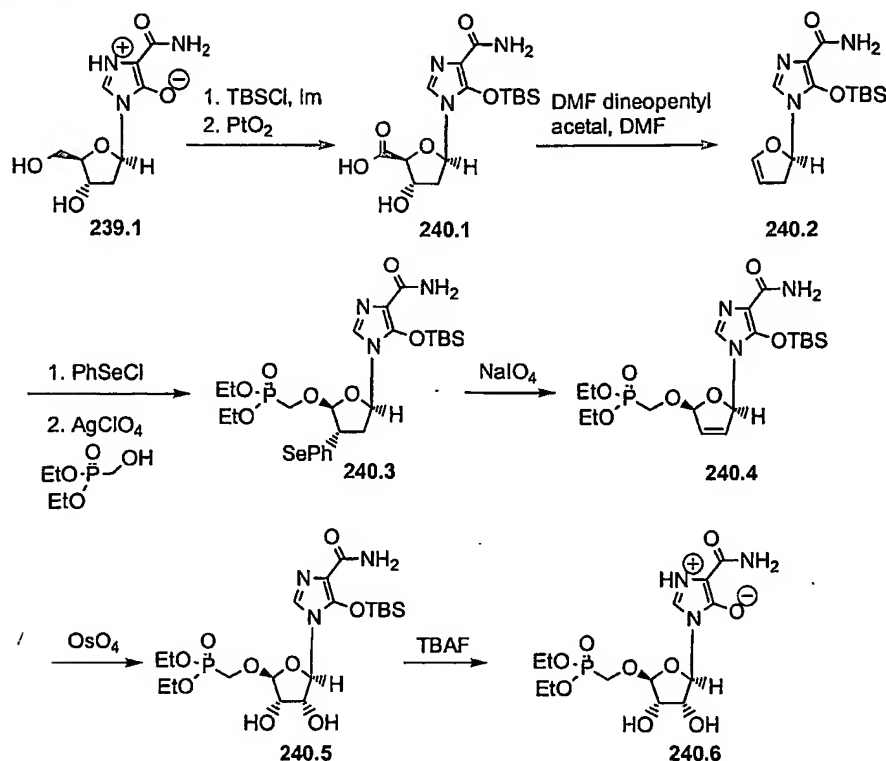
Example 239



The preparation of the phosphonate esters is illustrated above.

Compound **239.1**, 5-hydroxy-1-(4-hydroxy-5-hydroxymethyl-tetrahydro-furan-2-ylmethyl)-1H-imidazole-4-carboxylic acid amide, prepared by addition of the imidazole base (JP Kokai 76 88965) onto the 3,5-bis-protected 2-deoxy-D-*erythro*-pentofuranosyl chloride (Hayashi, M. et al., *Chem. Pharm. Bull.*, **1975**, *23*, 1, 245; Montgomery, J. A. et al., *J. Med. Chem.*, **1969**, *12*, 3, 498; and Iwamoto, R. H. et al., *J. Med. Chem.*, **1963**, *6*, 684), is protected on the imidazole-4-ol. Oxidation of the 5'-OH followed by elimination provides glycal, **239.2**. (See the procedure in Zemlicka J. et al., *J. Am. Chem. Soc.*, **1972**, *94*, 9, 3213.) Selenoetherification provides the protected phosphonate, **239.3** (Kim, C. et al., *J. Org. Chem.*, **1991**, *56*, 2642). Oxidative elimination of the phenylselenide (as described in Kim, C. et al., *J. Org. Chem.*, **1991**, *56*, 2642) followed by stereoselective dihydroxylation provides the diol **239.4**. Finally, the protecting group is removed to provide compound **239.5**.

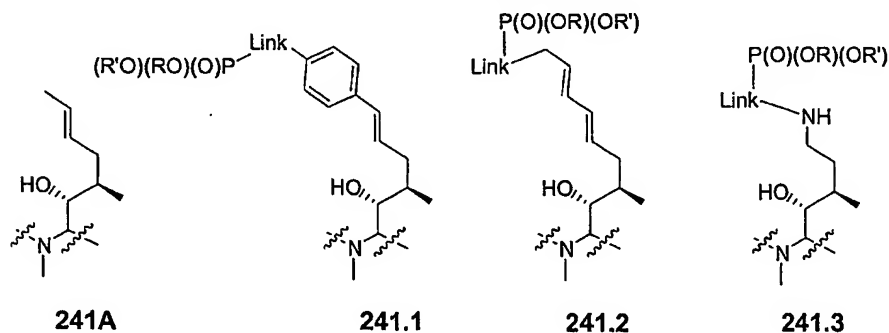
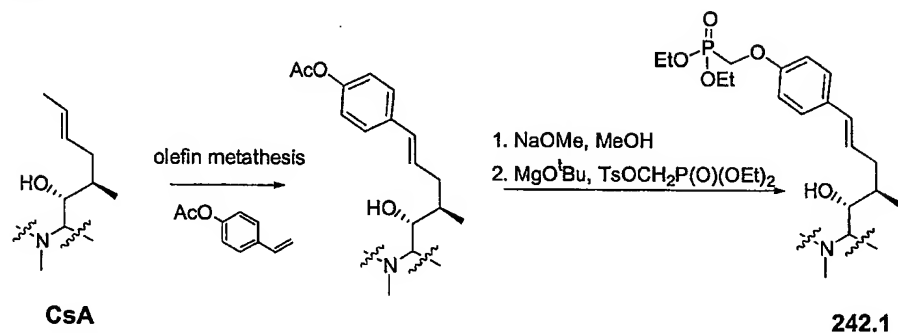
Example 240



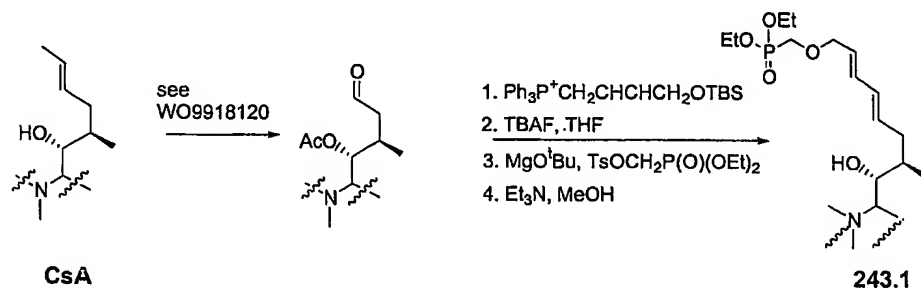
The preparation of the phosphonates is illustrated above. Specifically, compound **239.1**, 5-hydroxy-1-(4-hydroxy-5-hydroxymethyl-tetrahydrofuran-2-ylmethyl)-1H-imidazole-4-carboxylic acid amide, is first protected using a TBS group. Subsequent oxidation with PtO_2 proceeds to provide carboxylic acid **240.1**. Decarboxylative elimination is achieved using dimethylformamide dineopentyl acetal in DMF at high temperature (Zemlicka J. et al., *J. Am. Chem. Soc.*, 1972, 94, 9, 3213). Once the furanoid glycal **240.2** is in hand, it is treated with silver perchlorate in the presence of diethyl(hydroxymethyl)phosphonate (Phillion, D. et al., *Tetrahedron Lett.*, 1986, 27, 1477) to provide the phosphonate **240.3** (Kim, C. et al., *J. Org. Chem.*, 1991, 56, 2642). Oxidative elimination of the selenide followed by dihydroxylation using osmium tetroxide provides a diol, **240.5**, with the desired stereochemistry. Deprotection of the TBS group can be achieved using TBAF to provide compound **240.6**.

Example 241

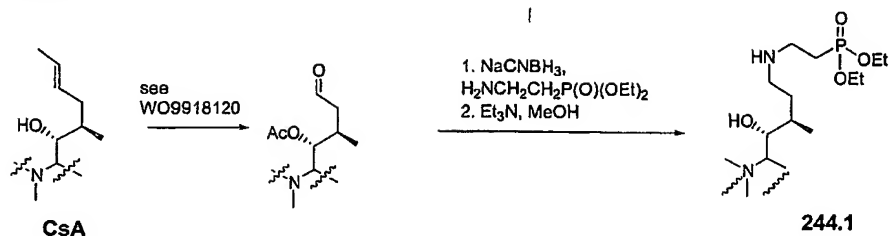
The synthesis of compounds of the invention that are analogs of CsA, 241A, (and related cyclosporins) are shown herein.

**Example 242**

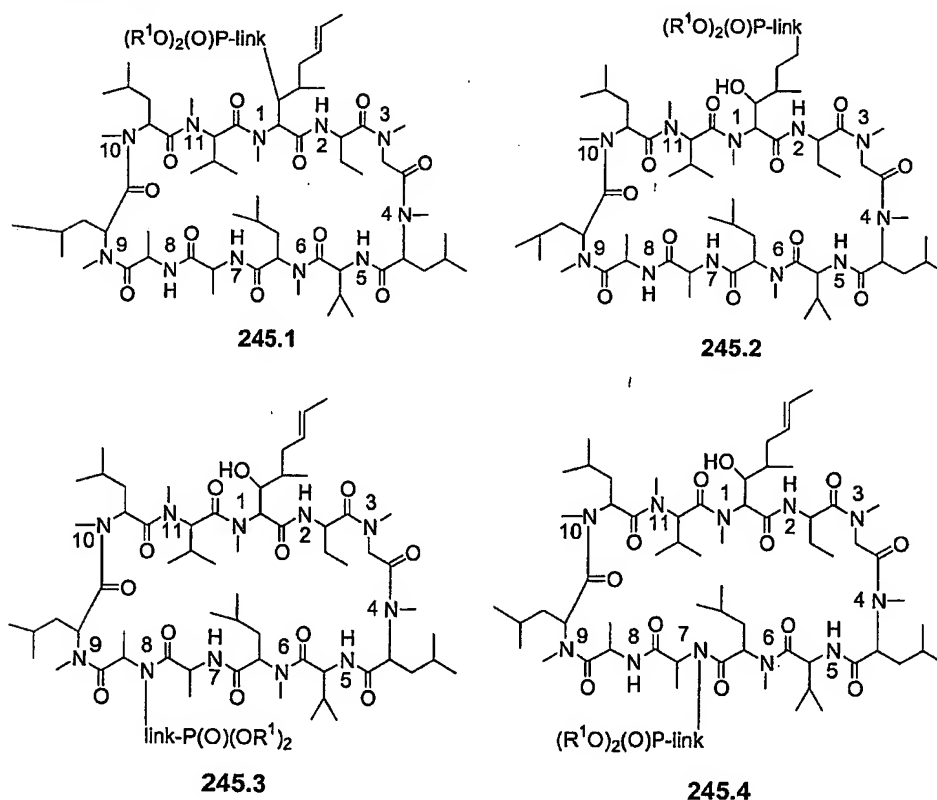
The syntheses of phosphonate compounds of the invention and of intermediate compounds necessary for their synthesis are illustrated herein. The olefin metathesis methodology is described in *J. Med. Chem.*, 2003, 46, 674.

Example 243

The preparation of compounds of the invention having phosphonate groups and intermediate compounds useful for their synthesis are illustrated herein.

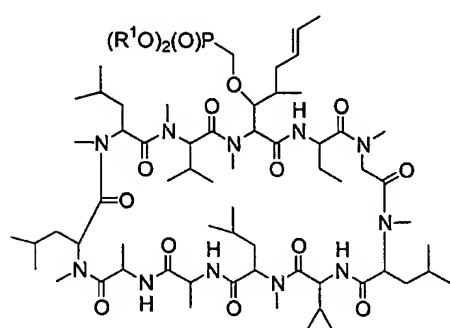
Example 244

The preparation of compounds of the invention having phosphonate groups and intermediate compounds useful for their synthesis are illustrated herein.

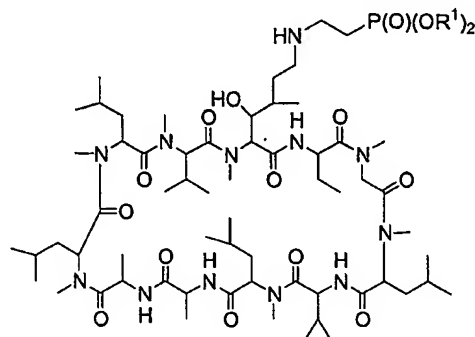
Example 245

The conversion of various substituents into the group link-P(O)(OR¹)₂, where R¹ is as defined above, or indeed the final stage of P(O)RR⁰, as defined above, can be effected at any convenient stage of the synthetic sequence, or in the final step. The selection of an appropriate step for the introduction of the

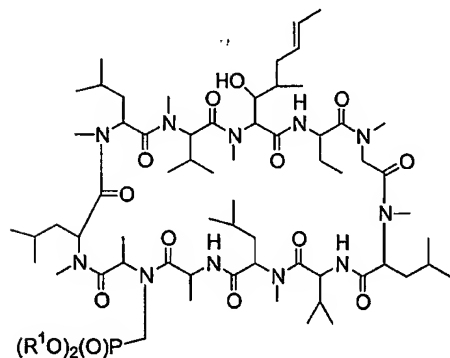
phosphonate substituent is made after consideration of the chemical procedures required and the stability of the substrates to those procedures. It may be necessary to protect reactive groups, for example hydroxyl, amino, during the introduction of the group link-P(O)(OR¹)₂ or P(O)RR^o. Specific examples of compounds of the invention having formulae, 245.5 – 245.8, are shown herein.



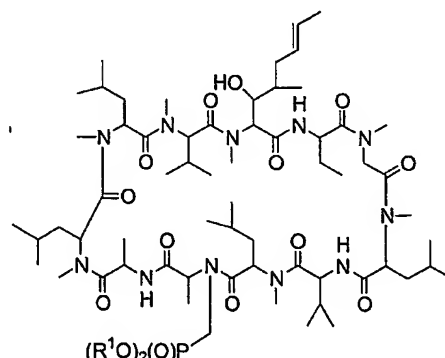
245.5



245.6



245.7

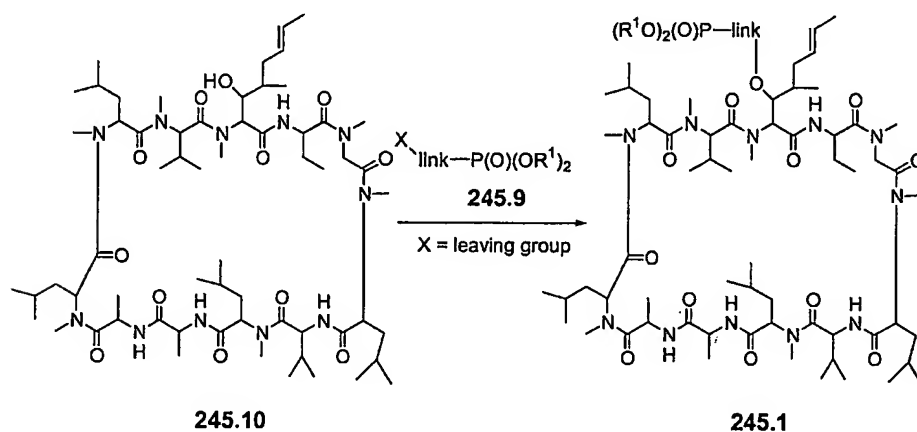


245.8

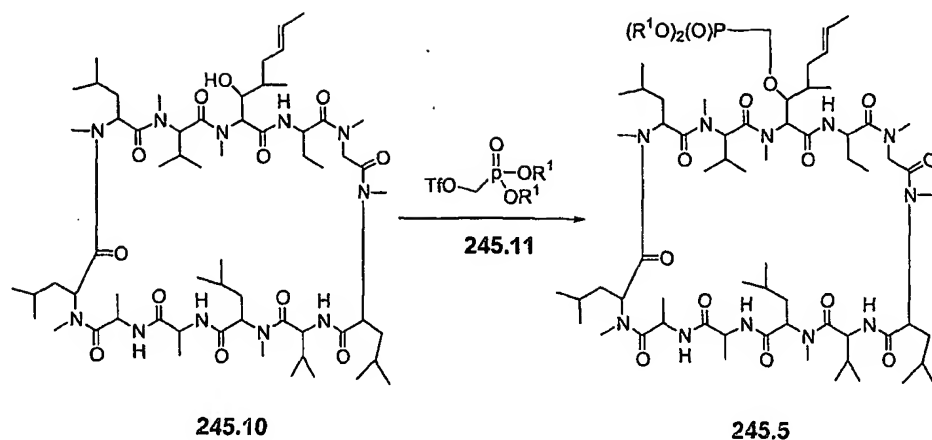
The compounds of the invention are synthesized according to the methods described herein. Exemplary intermediate phosphonate esters are shown, wherein R¹ is hydrogen, alkyl, aryl, haloalkyl, alkenyl, aralkyl, or aryl. These compounds can be used to prepare the compounds of the invention, such as those illustrated herein, by one skilled in the art, using known methods for synthesis of substituted phosphonates. These methods are similar to those described for the synthesis of amides. The preparation of amides from carboxylic acids and derivatives is described, for example, in "Organic Functional Group Preparations," by S.R. Sandler and W. Karo, Academic Press, 1968, p. 274. Further methods described in the following Examples for the

synthesis of the phosphonate diesters and can in some cases be applied to the synthesis of phosphoramides.

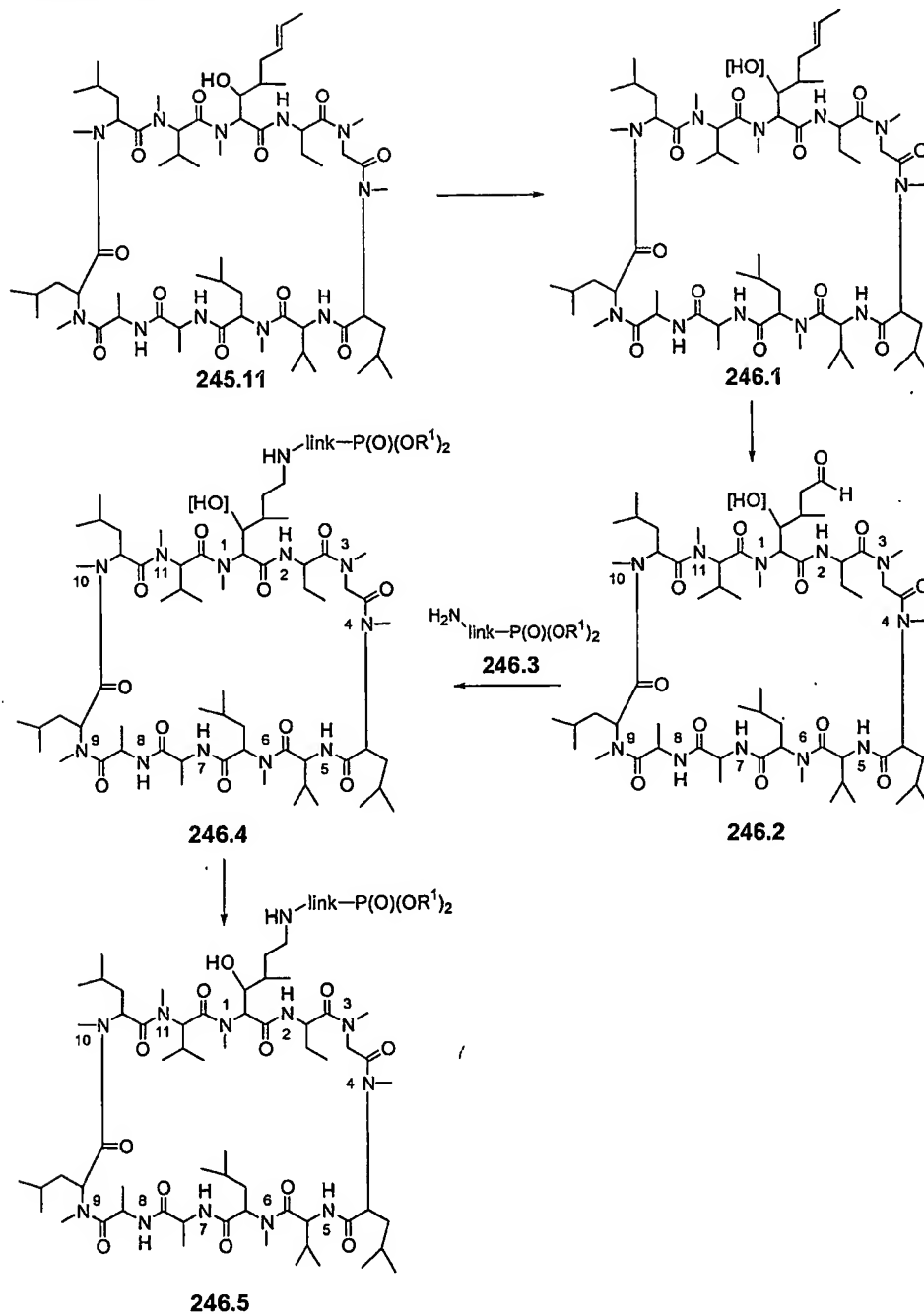
The phosphonate esters, **245.1-245.4**, for conversion into the phosphonate moieties bearing an amino acid, or a lactate esters are shown in herein. Cyclosporin A (CsA) **245.10**, can be purchased from Sigma Aldrich, synthesized (See U.S. Patent No. 4,396,542) or obtained from biological sources as described in U.S. Patent No. 4,117,118. Other cyclosporin derivatives can be either synthetic in nature (See U.S. Patent No. 4,396,542) or isolated by similar means to CsA (See U.S. Patent No. 6,410,696 B1).



The preparation of the phosphonate linkage to CsA through the hydroxyl group of amino acid 1 to give compounds of formula **245.1** is illustrated herein. CsA, **245.10**, is dissolved in a suitable solvent such as, for example, DMF or other non-protic solvent, and is then treated with the phosphonate reagent, **245.9**, bearing a leaving group, for example, bromine, mesyl, tosyl, or trifluoromethanesulfonyl in the presence of a suitable organic or inorganic base.



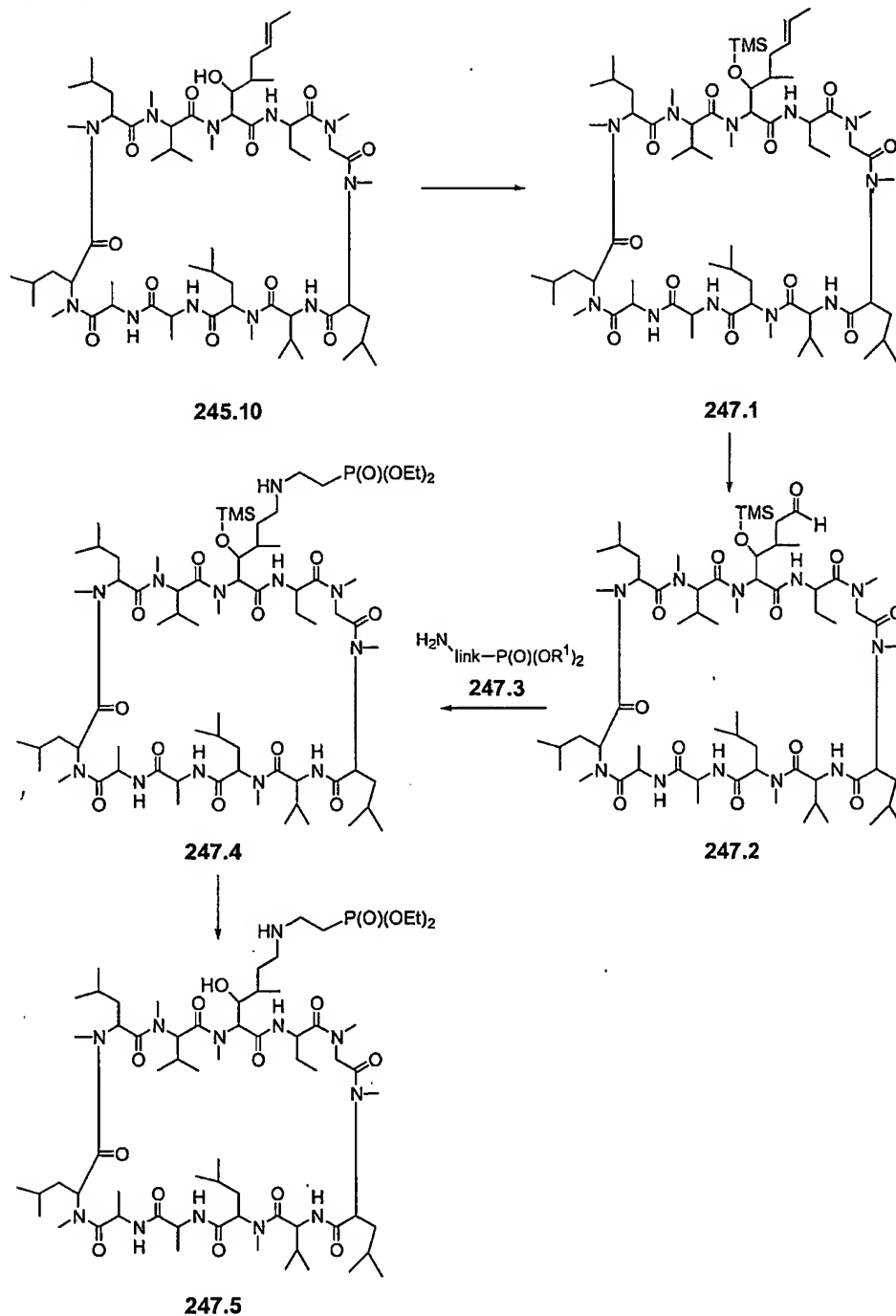
For example, **245.10** dissolved in DMF, is treated with one equivalent of sodium hydride and one equivalent of (toluene-4-sulfonylmethyl)-phosphonic acid dibenzyl ester **245.11**, prepared according to the procedures in in *J. Org. Chem.* **1996**, *61*, 7697, to give CsA phosphonate **245.5**. Using the above procedure but employing different phosphonate reagents, **245.9**, in place of compound, **245.11**, there are obtained the corresponding products, **245.1**, bearing different linking groups.

Example 246

The preparation of CsA - phosphonate conjugates is illustrated above. The hydroxyl group of amino acid 1 is first protected with a suitable protecting group, for example silyl ethers, benzyl ethers, trityl ethers etc as described in Greene and Wuts, "Protecting Groups in Organic Synthesis," 3rd Edition, John Wiley and Sons, Inc. The protected product, **246.1**, is then treated with an

oxidizing agent, many examples of which are described in "Comprehensive Organic Transformations," John Wiley & Sons, 2nd Ed, R. C. Larock, p 1211-1215 to give the aldehyde, 246.2.

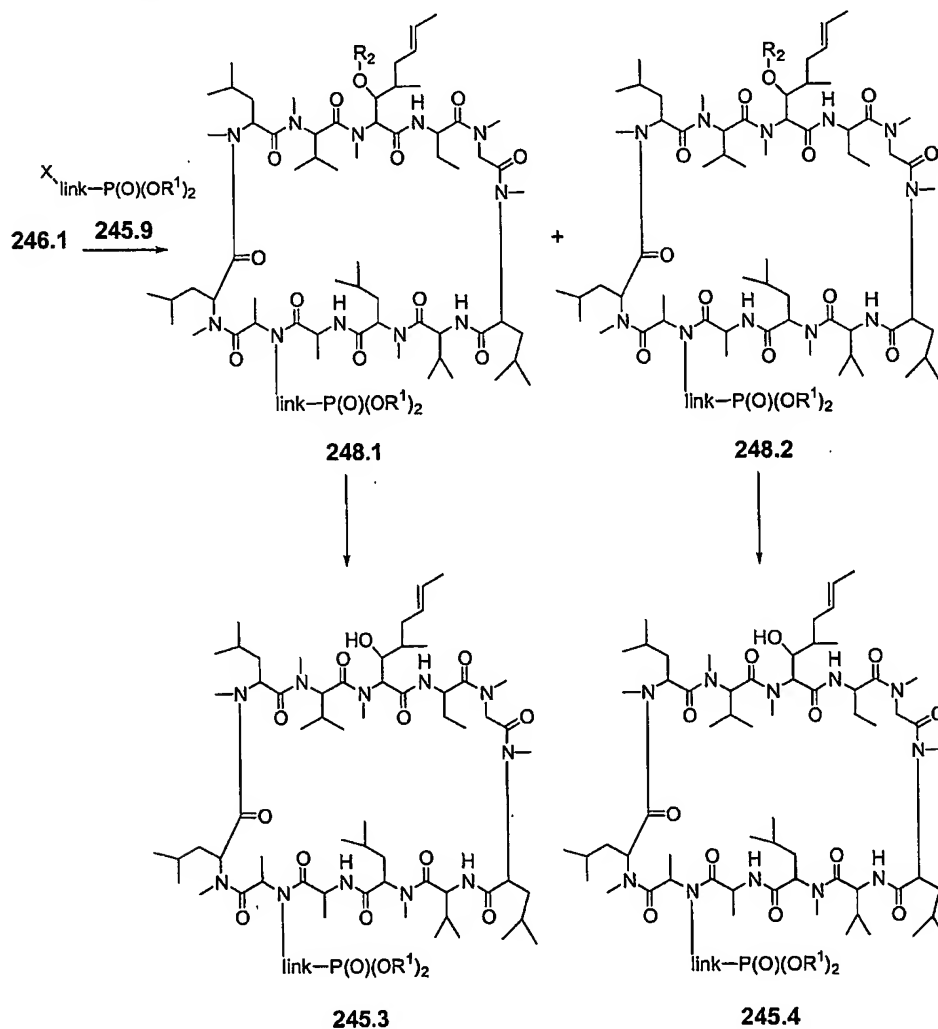
Aldehyde, 246.2, is then treated with a amine phosphonic acid ester of the general formula 246.3 under reductive amination conditions to afford amine, 246.4. The preparation of amines by means of reductive amination procedures is described, for example, in "Comprehensive Organic Transformations," by R. C. Larock, 2nd edition, p. 835. In this procedure, the amine component and the aldehyde component are reacted together in the presence of a reducing agent such as, for example, borane, sodium cyanoborohydride or diisobutylaluminum hydride, to yield the amine product. Finally, deprotection of the hydroxyl group following procedures documented in Greene and Wuts, "Protecting Groups in Organic Synthesis," 3rd Edition, John Wiley and Sons Inc. p116-121 gives the phosphonate 246.5.

Example 247

For example, **245.10**, is treated in pyridine and dichloromethane with trimethylsilyl chloride, as described in U.S. Patent No. 6,410,696 B1, to give silyl ether, **247.1**. Silyl ether, **247.1**, is then treated with ozone followed by work up with dimethyl sulfide to give aldehyde **247.2**. Aldehyde, **247.2**, is treated with one equivalent of the hydrochloride salt of (2-amino-ethyl)-phosphonic acid

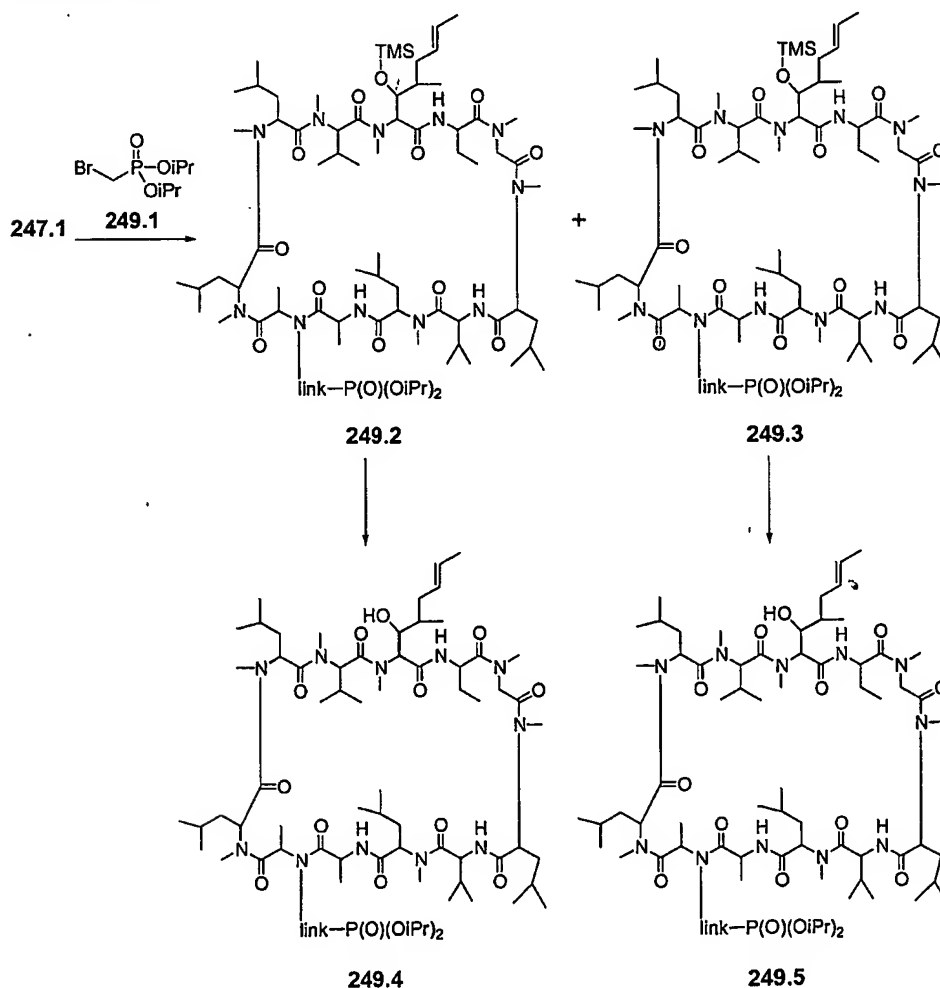
ester diethyl ester, **247.3**, prepared according to *J. Med. Chem.*, **1998**, *41*, 23, p4439, and a suitable base, e.g. hunigs base, triethylamine, or the likes, until the imine is formed. The intermediate imine solution is then treated with sodium cyanoborohydride to give the amine, **247.4**. Amine, **247.4**, is then deprotected by treatment with TBAF in an aprotic solvent such as THF or dioxane to give phosphonate, **247.5**. Using the above procedure but employing different phosphonate reagents, **246.3**, in place of phosphonate, **247.3**, there are obtained the corresponding products, **246.5**, bearing different linking groups.

Example 248



The preparation of CsA phosphonate conjugates where the phosphonate is linked onto the alanine nitrogen in amino acids 7 and 8, compounds, **245.3** and **245.4**, is illustrated herein. Protected CsA, **246.1** (Example 246), is treated with a base, sufficiently basic to remove the amide proton, for example, metal hydrides, metal amides. The product is then treated with a phosphonate reagent, **245.9**, bearing a leaving group such as, for example, bromine, mesyl, tosyl, or trifluoromethanesulfonyl phosphonates, to give **248.1** and **248.2**. The alkylated products are then separated by chromatography and independently deprotected using conventional conditions described in Greene and Wuts, "Protecting Groups in Organic Synthesis," 3rd Edition, John Wiley and Sons Inc. p116-121 to give compounds having Formulae, **245.3** and **245.4**.

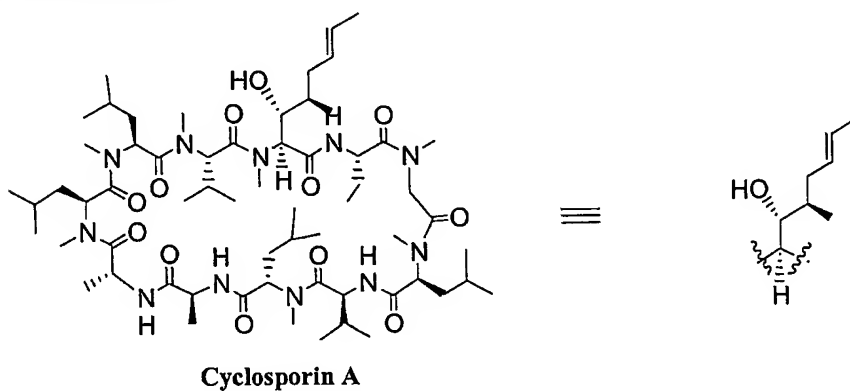
Example 249

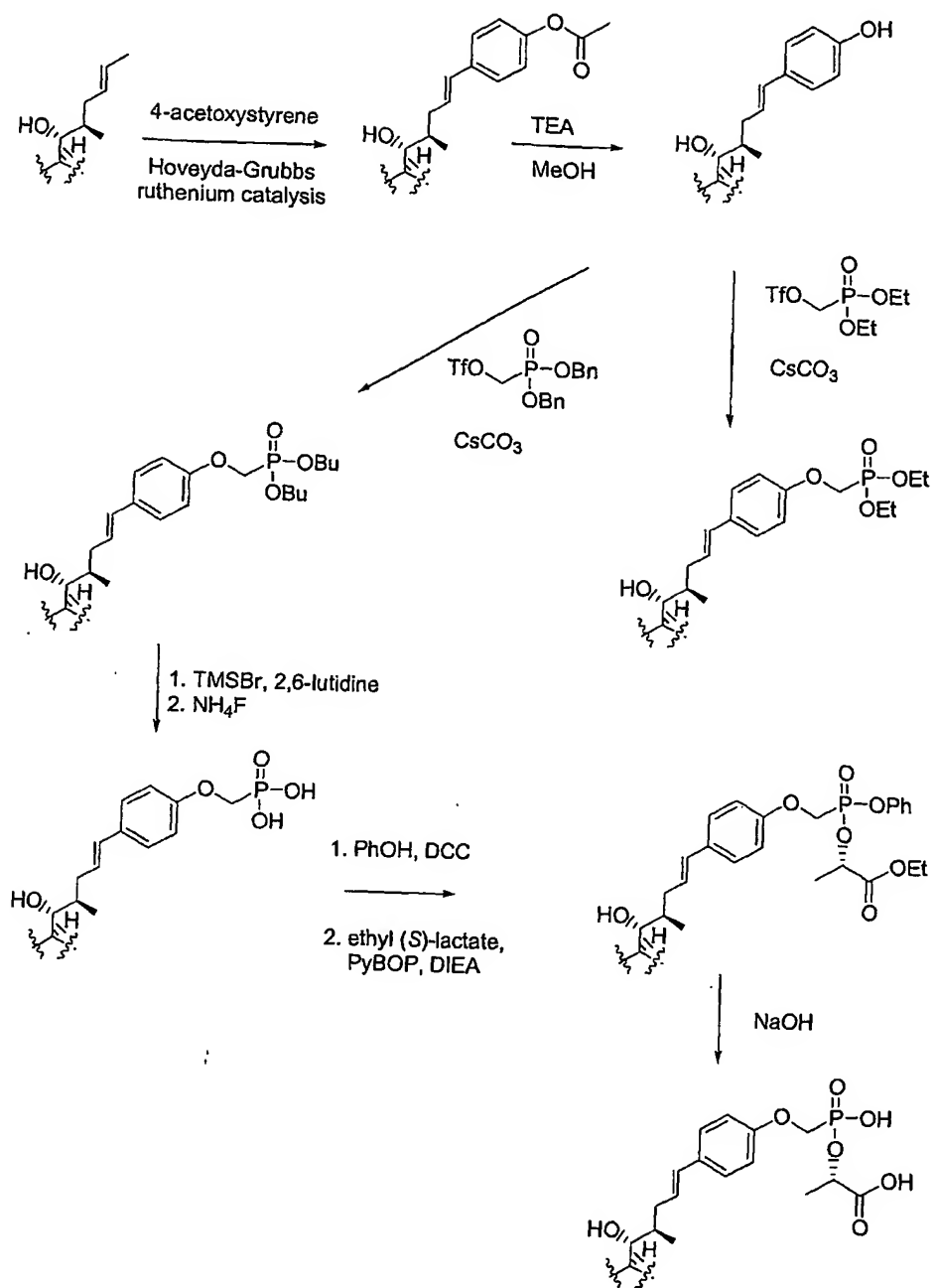


Silyl ether, **247.1**, in toluene is treated with sodium hydride and 15-crown-5-ether followed by one equivalent of bromomethyl phosphonic acid diallyl ester, **249.1** (Lancaster), to give phosphonates, **249.2** and **249.3**, respectively. Phosphonates, **249.2** and **249.3**, are deprotected by treatment with TBAF in an aprotic solvent such as THF or dioxane to give compounds having Formulae, **249.4** and **249.5**, respectively, wherein the linkage is a methylene group. Using the above procedure but employing different phosphonate reagents, **245.9**, in place of phosphonate reagents, **249.1**, there are obtained the corresponding products having Formulae, **245.3** and **245.4**, with different linking groups.

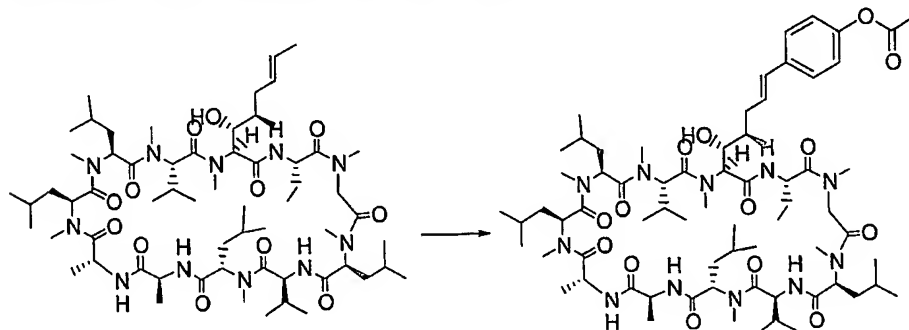
Example 250

The preparation of compounds of the invention having phosphonate groups and intermediate compounds useful for their synthesis are illustrated herein. The substructure on the right is meant to represent Cyclosporin A in the following examples.





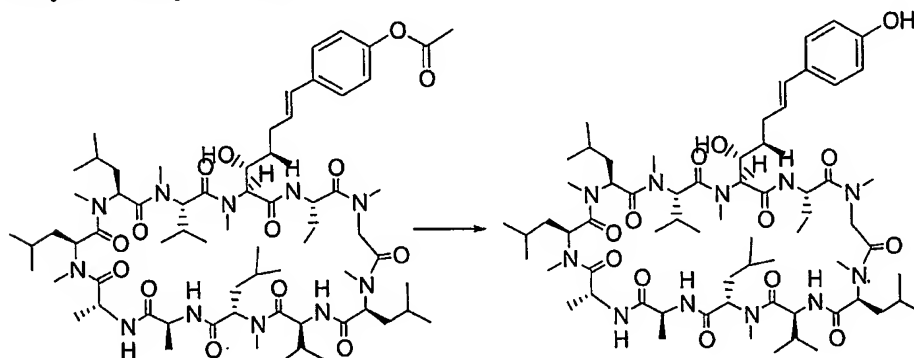
Example 251: *cyclo-[[[(2S, 3R, 4R, 6E)-7-(4-Acetoxyphenyl)-4-methyl-3-hydroxy-2-(methylamino)-6-heptenoyl]-L-2-aminobutyryl-sarcosyl-N-methyl-L-leucyl-L-valyl-N-methyl-L-leucyl-L-alanyl-D-alanyl-N-methyl-L-leucyl-N-methyl-L-leucyl-N-methyl-L-valyl]*



A mixture of cyclosporin A (360 mg, 0.3 mmol), 4-acetoxystyrene (730 mg, 4.5 mmol) and (1,3-bis-(2,4,6-trimethylphenyl)-2-imidazolidene)dichloro-(*O*-isopropoxyphenylmethylene)ruthenium (Hoveyda-Grubbs catalyst, 20 mg, 0.032 mmol) in dichloromethane (1 mL) was purged with nitrogen and stirred under reflux for 16 hours. After cooling, the reaction mixture was purified by silica gel column chromatography using MeOH - CH₂Cl₂ to provide the product as a solid (395 mg, 99 %).

MS (*m/z*) 1322.9 [M+H]⁺, 1344.9 [M+Na]⁺; HPLC retention time 3.3 min. (relative to 4.1 min. of cyclosporin A; Phenominex Synergi 4 micron hydro-RP 80A 50x4.6 mm; solvents, 35 % water and 65% acetonitrile; flow rate 2 mL/min.; column temperature 60 °C).

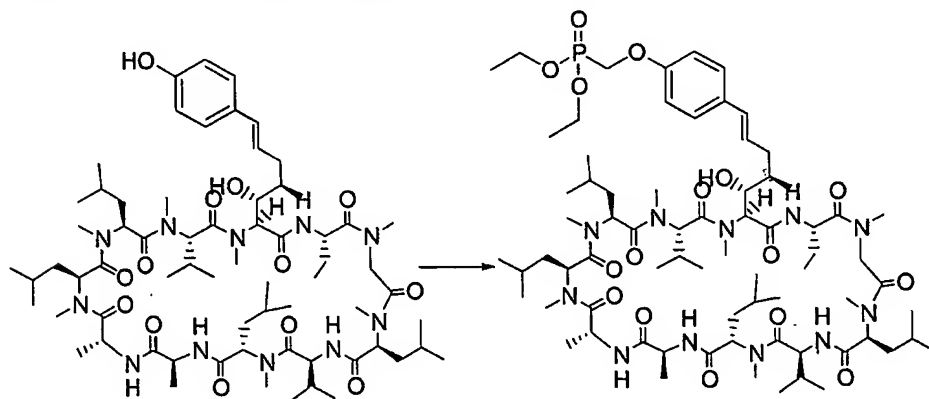
Example 252 *cyclo-[[[(2S, 3R, 4R, 6E)-7-(4-Hydroxyphenyl)-4-methyl-3-hydroxy-2-(methylamino)-6-heptenoyl]-L-2-aminobutyryl-sarcosyl-N-methyl-L-leucyl-L-valyl-N-methyl-L-leucyl-L-alanyl-D-alanyl-N-methyl-L-leucyl-N-methyl-L-leucyl-N-methyl-L-valyl]*



A solution of *cyclo*-[[*(2S, 3R, 4R, 6E)*-7-(4-acetoxyphenyl)-4-methyl-3-hydroxy-2-(methylamino)-6-heptenoyl]-L-2-aminobutyryl-sarcosyl-*N*-methyl-L-leucyl-L-valyl-*N*-methyl-L-leucyl-L-alanyl-D-alanyl-*N*-methyl-L-leucyl-*N*-methyl-L-leucyl-*N*-methyl-L-valyl] (385 mg, 0.29 mmol) and triethylamine (1 mL) in MeOH (10 mL) was stirred at ambient temperature for 16 hours. The reaction mixture was concentrated *in vacuo* and the residue was purified by silica gel column chromatography using MeOH - CH₂Cl₂ to provide the desired product (310 mg, 83%).

MS (*m/z*) 1280.9 [M+H]⁺, 1278.8 [M-H]⁻; HPLC retention time 1.6 min. (relative to 4.0 min. of cyclosporin A; Phenomenex Synergi 4 micron hydro-RP 80A 50x4.6 mm; solvents, 35 % water and 65% acetonitrile; flow rate 2 mL/min.; column temperature 60 °C).

Example 253: *cyclo*-[[*(2S, 3R, 4R, 6E)*-7-(4-(Diethoxyphosphorylmethoxy)phenyl)-4-methyl-3-hydroxy-2-(methylamino)-6-heptenoyl]-L-2-amino-butyryl-sarcosyl-*N*-methyl-L-leucyl-L-valyl-*N*-methyl-L-leucyl-L-alanyl-D-alanyl-*N*-methyl-L-leucyl-*N*-methyl-L-leucyl-*N*-methyl-L-valyl]

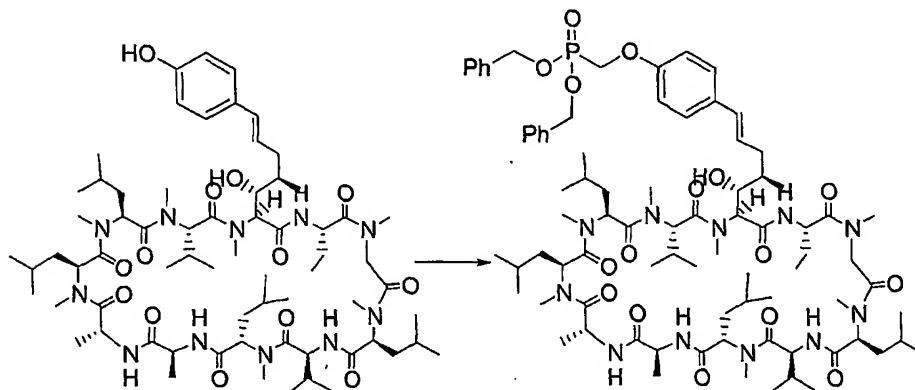


To a mixture of *cyclo*-[[*(2S, 3R, 4R, 6E)*-7-(4-hydroxyphenyl)-4-methyl-3-hydroxy-2-(methylamino)-6-heptenoyl]-L-2-aminobutyryl-sarcosyl-*N*-methyl-L-leucyl-L-valyl-*N*-methyl-L-leucyl-L-alanyl-D-alanyl-*N*-methyl-L-leucyl-*N*-methyl-L-leucyl-*N*-methyl-L-valyl] (113 mg, 0.088 mmol) and cesium carbonate (33 mg, 0.1 mmol) in DMF (1 mL) was added trifluoromethanesulfonic acid diethoxyphosphorylmethyl ester (60 mg, 0.2 mmol). The mixture was stirred at room temperature for 16 hours. The reaction was quenched with 2 % aqueous lithium chloride and the mixture was extracted with ethyl acetate. The ethyl acetate extract was concentrated *in vacuo*. The residue was purified by silica gel

column chromatography to provide the desired product (310 mg, 83%) contaminated with the unreacted starting materials, which was further purified by RP HPLC using a Phenomenex Synergi 5 μ Hydro RP 80A column (50 x 21.2 mm) with eluents of H₂O - CH₃CN. The fractions containing the desired product were pooled and concentrated to dryness (62 mg, 49 %).

MS (*m/z*) 1431.0 [M+H]⁺, 1428.7 [M-H]⁻; ³¹P (121.4 MHz, CDCl₃) δ 19.5.

Example 254: *cyclo*-[[*(2S, 3R, 4R, 6E)*-7-(4-(Dibenzylphosphoryl-methoxy)phenyl)-4-methyl-3-hydroxy-2-(methylamino)-6-heptenoyl]-L-2-aminobutyryl-sarcosyl-N-methyl-L-leucyl-L-valyl-N-methyl-L-leucyl-L-alanyl-D-alanyl-N-methyl-L-leucyl-N-methyl-L-leucyl-N-methyl-L-valyl]

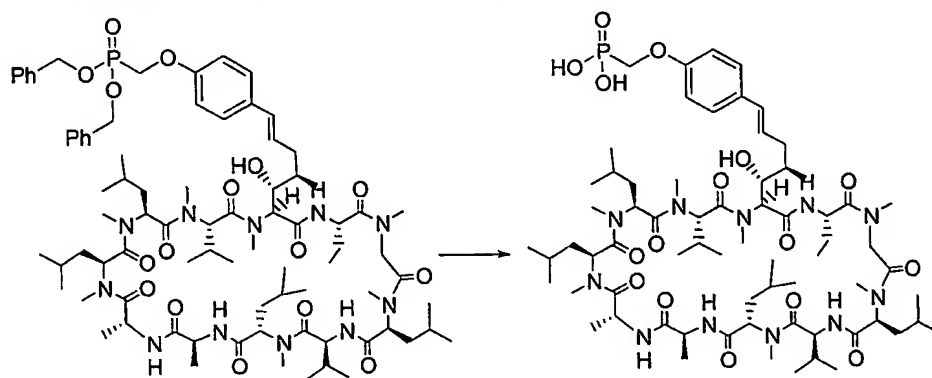


To a mixture of *cyclo*-[[*(2S, 3R, 4R, 6E)*-7-(4-hydroxyphenyl)-4-methyl-3-hydroxy-2-(methylamino)-6-heptenoyl]-L-2-aminobutyryl-sarcosyl-N-methyl-L-leucyl-L-valyl-N-methyl-L-leucyl-L-alanyl-D-alanyl-N-methyl-L-leucyl-N-methyl-L-leucyl-N-methyl-L-valyl] (300 mg, 0.234 mmol) and cesium carbonate (326 mg, 1 mmol) in DMF (2 mL) was added trifluoromethanesulfonic acid dibenzylphosphorylmethyl ester (60 mg, 0.2 mmol). The mixture was stirred at room temperature for 16 hours. The reaction mixture was filtered through Acrodisc (13 mm syringe filter with 0.45 micron Nylon membrane) and purified by RP HPLC using a Phenomenex Synergi 5 μ Hydro RP 80A column (50 x 21.2 mm) with eluents of H₂O- CH₃CN. The fractions containing the desired product were pooled and concentrated to dryness, affording a white solid (115 mg, 32 %).

MS (m/z) 1554.9 $[M+H]^+$, 1552.7 $[M-H]^-$; ^{31}P (121.4 MHz, CDCl_3) δ

20.5.

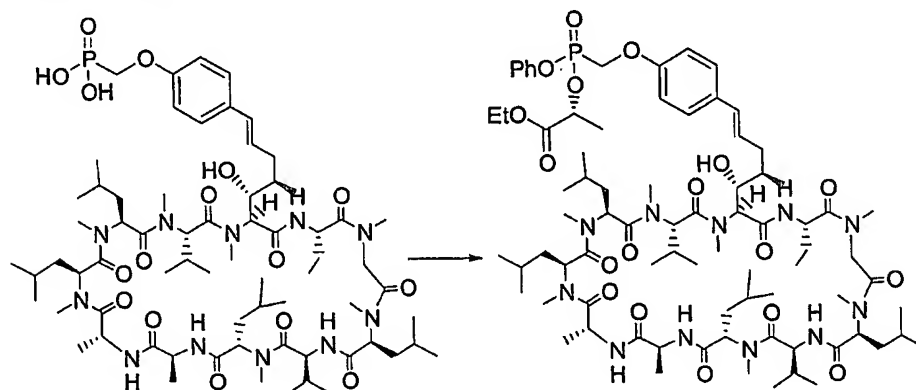
Example 255: *cyclo-[[[(2S, 3R, 4R, 6E)-7-(4-(Dihydroxyphosphoryl-methoxy)phenyl)-4-methyl-3-hydroxy-2-(methylamino)-6-heptenoyl]-L-2-aminobutyryl-sarcosyl-N-methyl-L-leucyl-L-valyl-N-methyl-L-leucyl-L-alanyl-D-alanyl-N-methyl-L-leucyl-N-methyl-L-leucyl-N-methyl-L-valyl]*



To a mixture of *cyclo-[[[(2S, 3R, 4R, 6E)-7-(4-(dibenzoyloxyphosphoryl-methoxy)phenyl)-4-methyl-3-hydroxy-2-(methylamino)-6-heptenoyl]-L-2-aminobutyryl-sarcosyl-N-methyl-L-leucyl-L-valyl-N-methyl-L-leucyl-L-alanyl-D-alanyl-N-methyl-L-leucyl-N-methyl-L-leucyl-N-methyl-L-valyl]* (115 mg, 0.074 mmol) and 2,6-lutidine (40 μL , 0.35 mmol) in dichloromethane (2 mL) was added trimethylsilyl bromide (50 μL , 0.35 mmol). The mixture was stirred at room temperature for 2 hours. The reaction was quenched with methanol (1 mL) and the mixture was concentrated. The residue was treated with a solution of ammonium fluoride (0.5 M, 2 mL), stirred for 1 hour, concentrated, and partitioned between dichloromethane and 1 N HCl. The dichloromethane layer was concentrated and the crude product was purified by RP HPLC using a Phenomenex Synergi 5 μ Hydro RP 80A column (50 x 21.2 mm) with eluents of 0.1 % TFA H_2O - 0.1 % TFA CH_3CN . The fractions containing the desired product were pooled and concentrated to dryness, affording a hygroscopic solid (68 mg, 63 %).

MS (m/z) 1374.9 $[M+H]^+$, 1373.1 $[M-H]^-$; HPLC retention time 0.3 min. (relative to 4.0 min. of cyclosporin A; Phenomenex Synergi 4 micron hydro-RP 80A 50x4.6 mm; solvents, 35 % water and 65% acetonitrile; flow rate 2 mL/min.; column temperature 60 $^\circ\text{C}$).

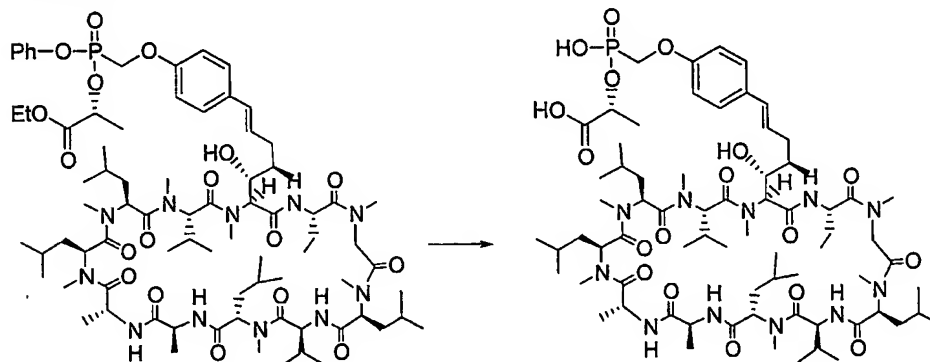
Example 256: *cyclo-[[[(2S, 3R, 4R, 6E)-7-(4-(1-(S)-Ethoxycarbonylethoxy)-phenoxyphosphorylmethoxy)phenyl)-4-methyl-3-hydroxy-2-(methylamino)-6-heptenoyl]-L-2-aminobutyryl-sarcosyl-N-methyl-L-leucyl-L-valyl-N-methyl-L-leucyl-L-alanyl-D-alanyl-N-methyl-L-leucyl-N-methyl-L-leucyl-N-methyl-L-valyl]*



A mixture of *cyclo-[[[(2S, 3R, 4R, 6E)-7-(4-(dihydroxyphosphoryl-methoxy)phenyl)-4-methyl-3-hydroxy-2-(methylamino)-6-heptenoyl]-L-2-aminobutyryl-sarcosyl-N-methyl-L-leucyl-L-valyl-N-methyl-L-leucyl-L-alanyl-D-alanyl-N-methyl-L-leucyl-N-methyl-L-leucyl-N-methyl-L-valyl]* (34 mg, 0.023 mmol), phenol (22 mg, 0.23 mmol), dicyclohexylcarbodiimide (47 mg, 0.23 mmol) and 4-(*N, N*-dimethylamino)pyridine (5.6 mg, 0.046 mmol) in DMF (2 mL) was stirred at 140 °C for 20 min. After cooling, the monophenyl monophosphonic acid product was purified by RP HPLC using a Phenomenex Synergi 5 μ Hydro RP 80A column (50 x 21.2 mm) with eluents of 0.1 % TFA H₂O - 0.1 % TFA CH₃CN. MS (*m/z*) 1450.9 [M+H]⁺, 1449.1 [M-H]⁻; ³¹P (121.4 MHz, CDCl₃) δ 14.9. This intermediate was mixed with ethyl (*S*)-(-)-lactate (40 mg, 0.34 mmol), PyBOP (80 mg, 0.15 mmol), diisopropylethylamine (45 μ L, 0.26 mmol) and DMF (1.7 mL). The resulting mixture was stirred at room temperature for 2 hours. After removal of insoluble impurities, the crude product was purified by RP HPLC using a Phenomenex Synergi 5 μ Hydro RP 80A column (50 x 21.2 mm) with eluents of 0.1 % TFA H₂O - 0.1 % TFA CH₃CN. The desired fractions were pooled and partitioned between acetonitrile and saturated aqueous sodium bicarbonate. The organic layer was concentrated to afford the product as a solid (12 mg, 34 %).

MS (m/z) 1573.1 $[M+Na]^+$, 1548.8 $[M-H]^-$; ^{31}P (121.4 MHz, $CDCl_3$) δ 15.3 and 17.4.

Example 257: *cyclo-[[[(2S, 3R, 4R, 6E)-7-(4-(1-(S)-Hydroxycarbonylethoxy)-hydroxyphosphorylmethoxy)phenyl)-4-methyl-3-hydroxy-2-(methylamino)-6-heptenoyl]-L-2-aminobutyryl-sarcosyl-N-methyl-L-leucyl-L-valyl-N-methyl-L-leucyl-L-alanyl-D-alanyl-N-methyl-L-leucyl-N-methyl-L-leucyl-N-methyl-L-leucyl-N-methyl-L-valyl]*

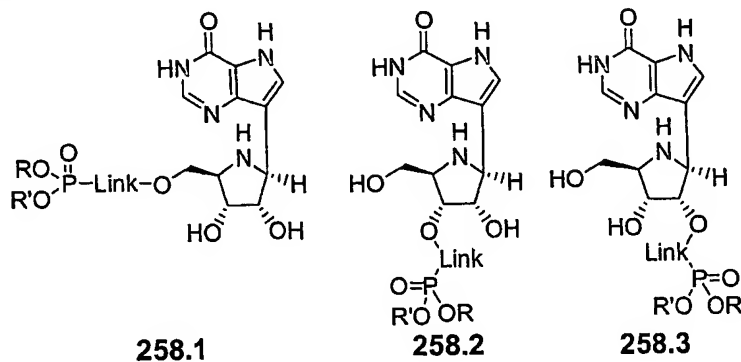


To a solution of *cyclo-[[[(2S, 3R, 4R, 6E)-7-(4-(1-(S)-ethoxycarbonylethoxy)phenoxyphosphorylmethoxy)phenyl)-4-methyl-3-hydroxy-2-(methylamino)-6-heptenoyl]-L-2-aminobutyryl-sarcosyl-N-methyl-L-leucyl-L-valyl-N-methyl-L-leucyl-L-alanyl-D-alanyl-N-methyl-L-leucyl-N-methyl-L-leucyl-N-methyl-L-leucyl-N-methyl-L-valyl]* (5 mg, 3.2 μ mol) in a mixed solvent of water and acetonitrile (0.5 mL and 4.5 mL) was added 1 N NaOH (40 μ L). The solution was stirred at room temperature for 2 hours. The resulting reaction mixture was concentrated and purified by RP HPLC using a Phenomenex Synergi 5 μ Hydro RP 80A column (50 x 21.2 mm) with eluents of 0.1 % TFA H_2O - 0.1 % TFA CH_3CN . The desired fraction was concentrated to dryness affording the product as a solid (1.5 mg, 32 %).

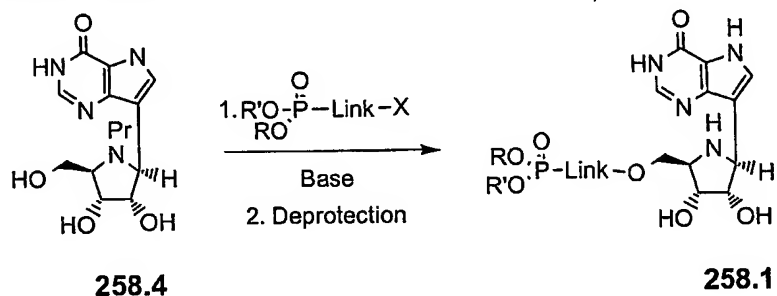
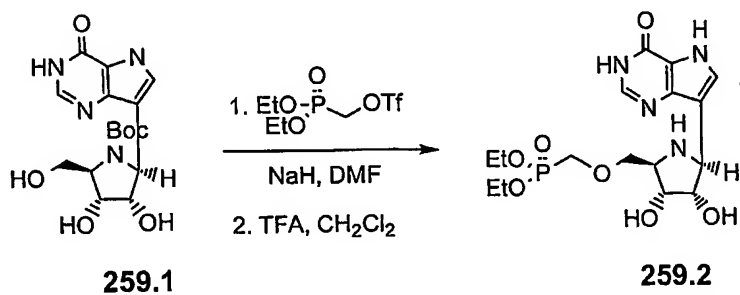
MS (m/z) 1446.9 $[M+H]^+$, 1444.9 $[M-H]^-$; HPLC retention time 0.2 min. (relative to 4.0 min. of cyclosporin A; Phenomenex Synergi 4 micron hydro-RP 80A 50x4.6 mm; solvents, 35 % water and 65% acetonitrile; flow rate 2 mL/min.; column temperature 60 $^{\circ}C$).

Example 258

The synthesis of compounds of the invention that are analogs of BCX-1777 is shown herein.



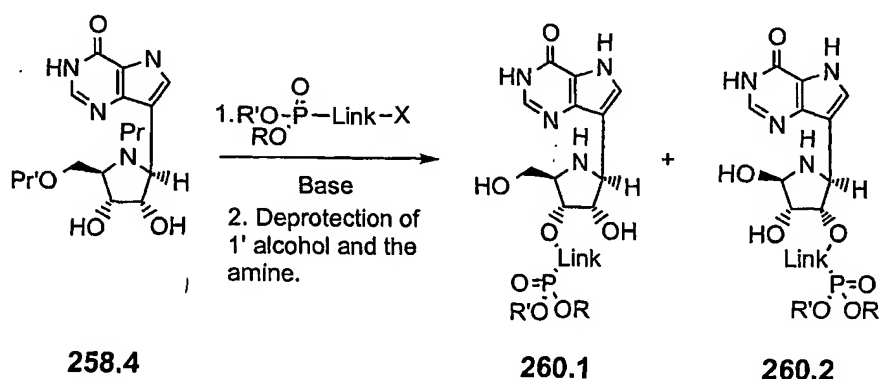
Compounds of the invention, such as, for example, **258.1** can be made according to the general route outlined herein.

**Example 259**

The Boc-protected (1S)-1-(9-deazaguanin-9-yl)-1,4-dideoxy-1,4-imino-D-ribose, compound, **259.1**, is prepared by stirring the (1S)-1-(9-deazaguanin-9-yl)-1,4-dideoxy-1,4-imino-D-ribose (WO 9,919,338 and Evans, G. B. et al., *Tetrahedron*, **2000**, *56*, 3053, also reported in Evans, G. B. et al., *J. Med. Chem.*

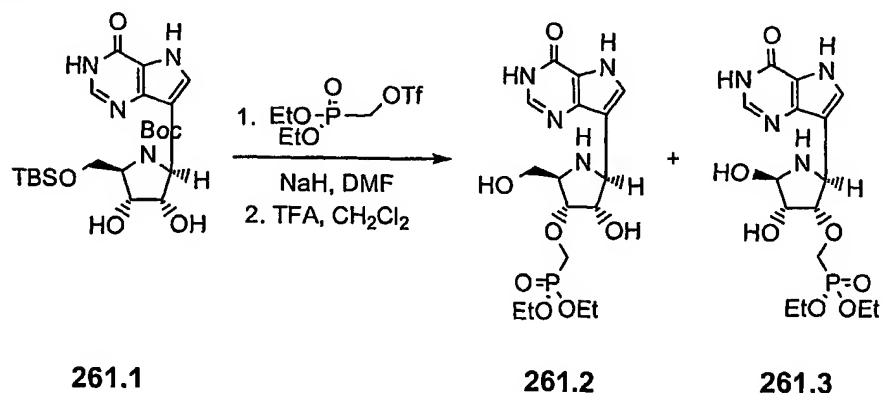
2003, 46, 3412) with BOC anhydride as described in Greene, T., "Protective Groups in Organic Synthesis," Wiley-Interscience, 1999. Compound, **259.1**, is then treated in a solvent such as tetrahydrofuran or dimethylformamide with a base such as sodium hydride. When bubbling ceases, diethyl phosphonomethyl-triflate (prepared according to *Tetrahedron Lett.*, 1986, 27, 1477) is added, yielding the desired phosphonate, **259.2**, after deprotection of the BOC group using trifluoroacetic acid (TFA).

Example 260



Compounds such as **260.1** and **260.2** can be made according to the general route outlined herein.

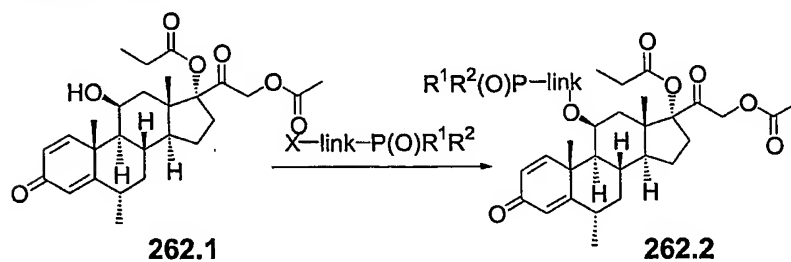
Example 261



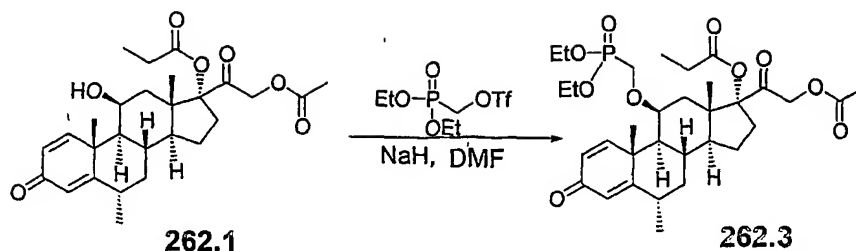
The Boc-protected (1S)-1-(9-deazaguanin-9-yl)-1,4-dideoxy-1,4-imino-D-ribose, compound, **259.1**, is prepared by stirring the (1S)-1-(9-deazaguanin-9-yl)-1,4-dideoxy-1,4-imino-D-ribose (WO 9,919,338 and Evans, G. B. et al., *Tetrahedron*, 2000, 56, 3053, also reported in Evans, G. B. et al., *J. Med. Chem.*

2003, 46, 3412) with BOC anhydride as described in Greene, T., "Protective Groups in Organic Synthesis," Wiley-Interscience, 1999. Subsequent protection of the primary alcohol using a TBS group can be achieved using TBSCl and imidazole in solvents such as CH_2Cl_2 as described in Greene, T., "Protective Groups in Organic Synthesis," Wiley-Interscience, 1999 to provide compound, 261.1. Compound, 261.1, is then treated in a solvent such as tetrahydrofuran or dimethylformamide with a base such as sodium hydride. When bubbling ceases, diethyl phosphonomethyltriflate (prepared according to *Tetrahedron Lett.*, 1986, 27, 1477) is added, yielding a mixture of the desired phosphonate diester, 261.2, and 261.3, after deprotection of the BOC group using trifluoroacetic acid (TFA). Compounds, 261.2, and 261.3, can be also prepared via a more complicated 2' OH protected analog of compound 261.1 followed by alkylation using the diethyl phosphonomethyltriflate to provide compound 261.2, exclusively. Compound 261.3 can also be prepared by installation of a different protecting group at the 3' OH position, followed by deprotection of 2' OH and alkylation with diethyl phosphonomethyltriflate at the 2' center followed by global deprotection.

Example 262

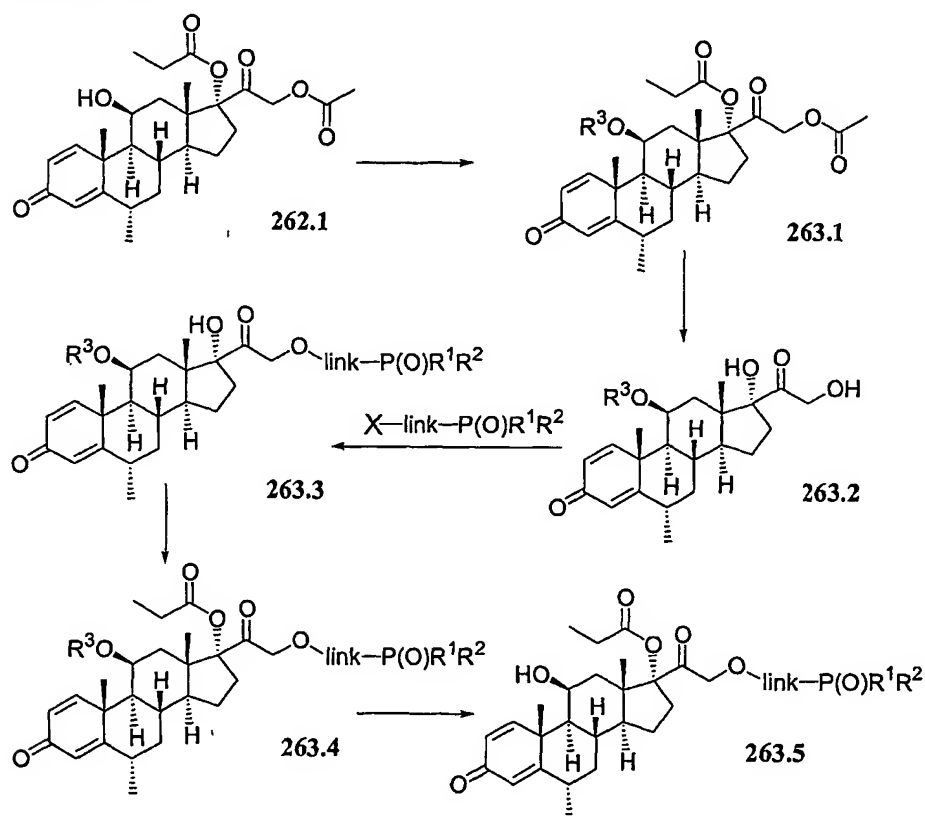


Representative compounds of the invention can be prepared as illustrated above. Derivatization at the C-11 hydroxy group is accomplished through alkylation of methylprednisolone aceponate 262.1 with the appropriate phosphonate, furnishing analogs of formula 262.2. A specific compound of the invention can be prepared as follows.



After sodium hydride extraction of the hydroxy proton in **262.1**, diethyl phosphonate triflate is added to afford ether **262.3**.

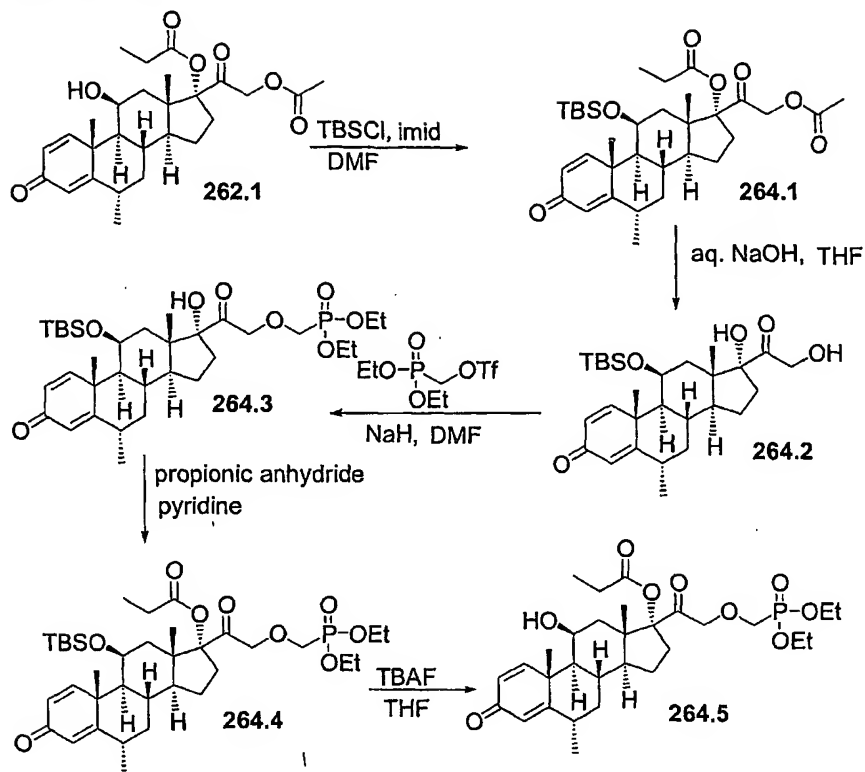
Example 263



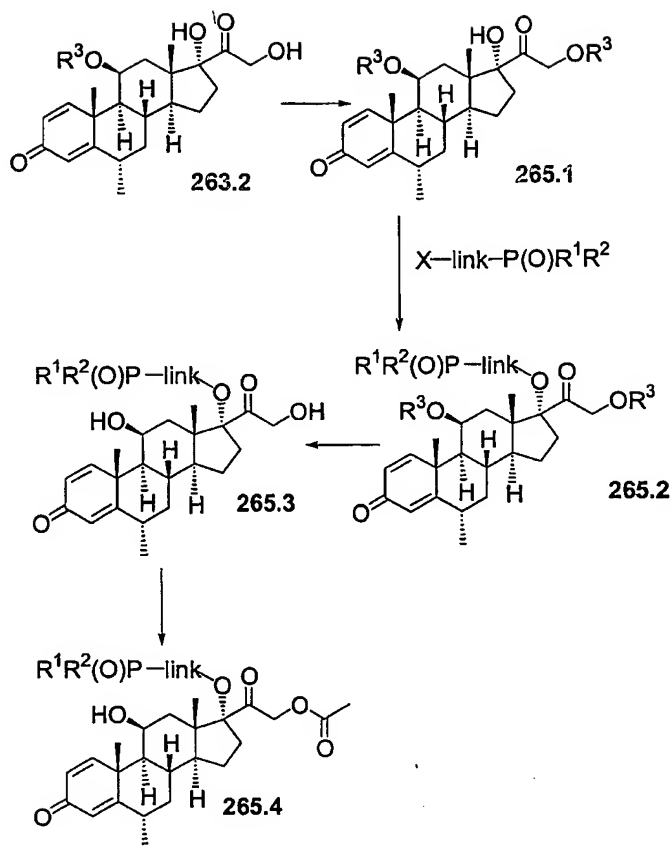
Representative compounds of the invention can be prepared as illustrated above by exploiting the reactivity differences among the three hydroxy groups available when methylprednisolone aceponate, **262.1**, is fully hydrolyzed. Following protection of the only exposed hydroxy group in **262.1**, intermediate, **263.1** is saponified to give diol, **263.2**. Alkylation at the primary hydroxy group with the appropriate phosphonate and subsequent acylation provides the

propionate ester, **263.4**. The desired product, **263.5**, is achieved after deprotection.

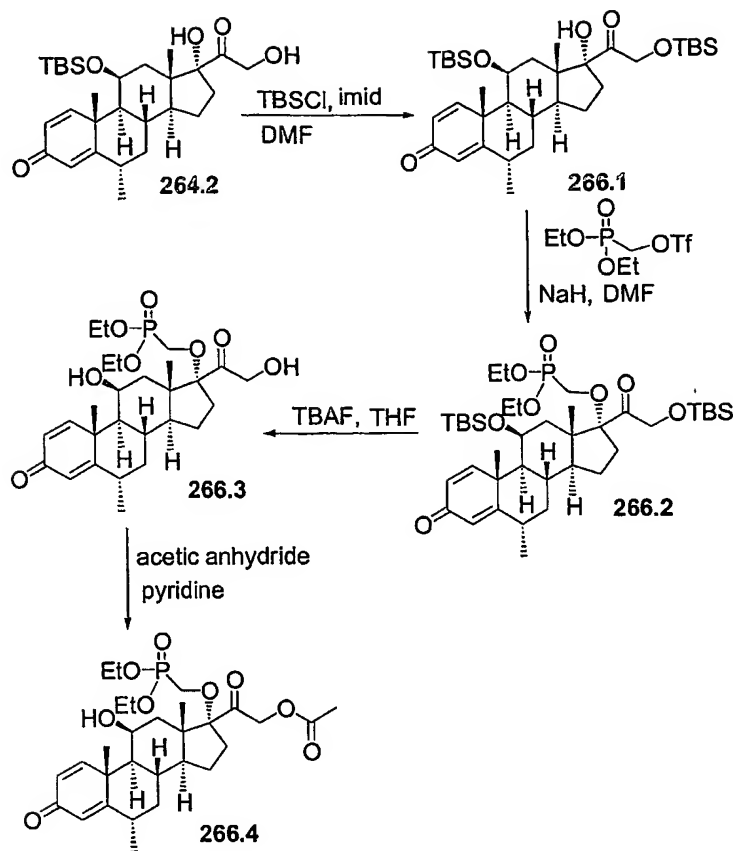
Example 264



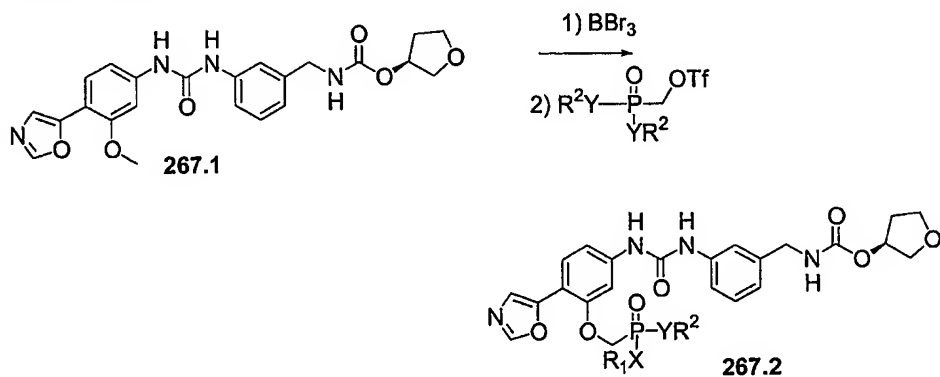
Methylprednisolone aceponate, **262.1**, is protected as its silyl ether using the standard TBSCl and imidazole conditions. (*J. Am. Chem. Soc.*, **1972**, *94*, 6190). Saponification of both ester moieties using aqueous sodium hydroxide provides the diol, **264.2**. The less sterically hindered primary hydroxy group is alkylated by the addition of sodium hydroxide and the phosphonate triflate. After treating intermediate, **264.3**, with propionic anhydride in pyridine, the previously hydrolyzed C-17 propionic ester is replaced. (*J. Med. Chem.*, **1980**, *23*, 430-437). TBAF deprotection of the silyl ether furnishes diethyl phosphonate **264.5**.

Example 265

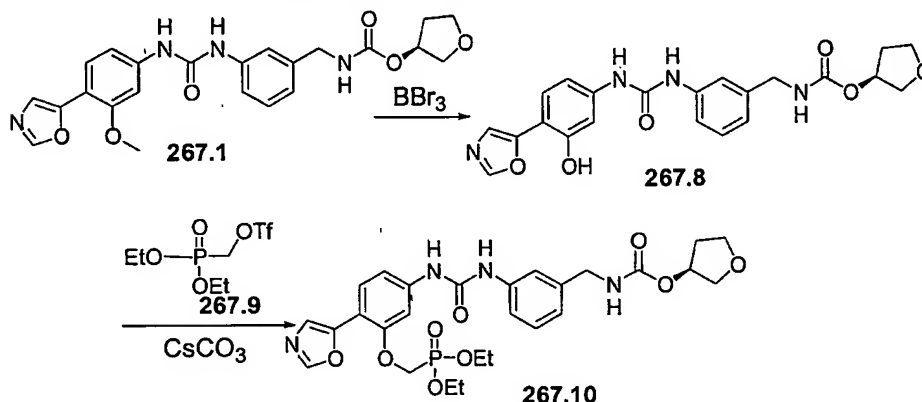
Representative compounds of the invention can be prepared as illustrated above. The two hydroxy groups of diol, 263.2, are regioselectively differentiated by protection at the primary site, thus allowing alkylation at the tertiary hydroxy group. The resulting phosphonate intermediate, 265.2, is then deprotected to afford the diol 265.3. Again the more accessible primary hydroxy group is acylated to produce the desired analog 265.4.

Example 266

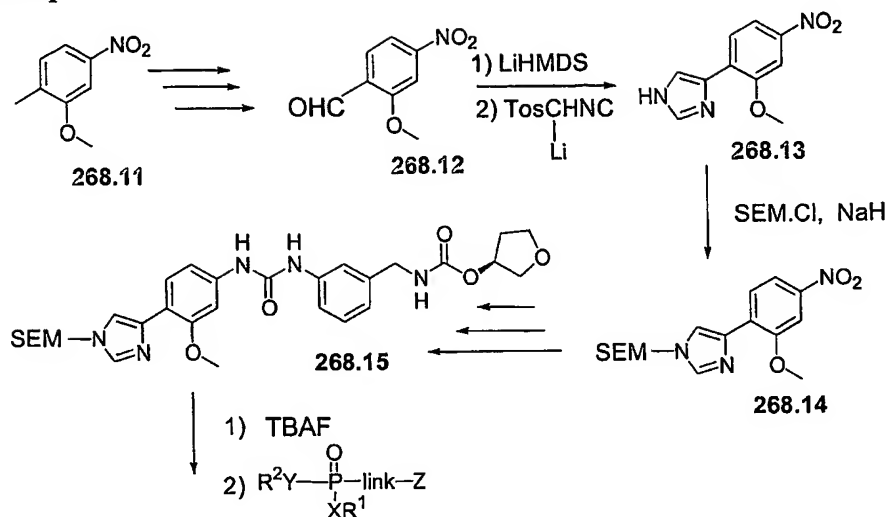
Diol, **264.2** (see example 264), is protected at the primary site as its silyl ether, **266.1**. Following alkylation with the diethyl phosphonate triflate, the resulting intermediate, **266.2**, is treated with TBAF to give diol, **266.3**. Acetic anhydride and pyridine are used to generate the final product **266.4**. (*J. Mol. Biol.*, **1972**, 72, 219).

Example 267

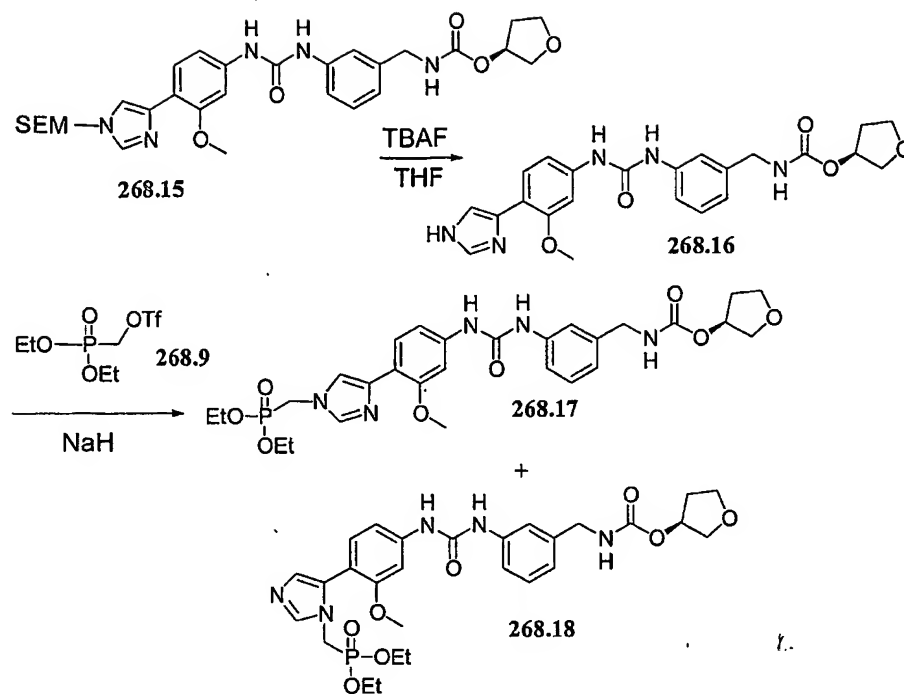
Representative compounds of the invention can be prepared as illustrated above. The phosphorus containing merimepodib analog **267.2** is synthesized from parent compounds by alkylation. Merimepodib **267.1** is obtained by the procedure as described in U.S. Patent No. 6,054,472 and U.S. Patent No. 6,344,465. The methoxy group of merimepodib **267.1** is demethylated to phenolic OH using a suitable reagent, such as boron tribromide. The phosphonate moiety is introduced to the phenolic OH in a suitable aprotic solvent such as, DMF and is then treated with the phosphonate reagent bearing a leaving group, for example, bromine, mesyl, tosyl, or trifluoromethanesulfonyl, in the presence of a suitable organic or inorganic base. A specific compound of the invention can be prepared as follows.



A solution of **267.1** in dichloromethane is treated with boron tribromide to obtain the demethylated compound **267.8**. Compound **267.8** is then treated with cesium carbonate and one equivalent of (trifluoromethanesulfonyloxy)-methylphosphonic acid diethyl ester **267.9** to give merimepodib-phosphonate **267.10**. Using the above procedure but employing different phosphonate reagents, the corresponding products **267.2** bearing different linking group can be obtained.

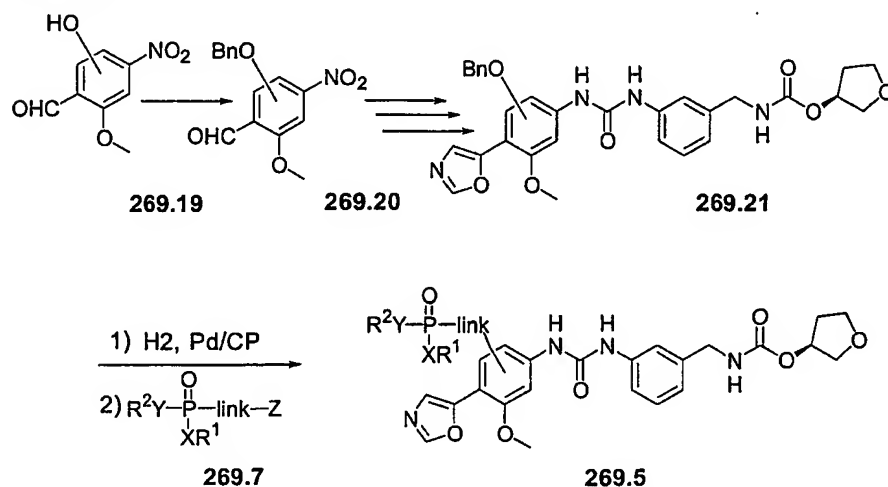
Example 268

Representative compounds of the invention can be prepared as illustrated above. The imidazole containing intermediate 268.13 is synthesized from an aldehyde 268.12 by the procedure of Shih in *Tetrahedron Lett.* 1993, 34, 595. Compound 268.12 is prepared by a two-step procedure described in U.S. Patent No. 5,807,876, U.S. Patent No. 6,054,472, and U.S. Patent No. 6,344,465. The imidazole is protected using suitable reagent, for example, 2-(trimethylsilyl)ethoxymethyl (SEM) chloride, and the compound 268.14 is converted to 268.15 by the similar procedure described for the synthesis of 197.1 in U.S. Patent No. 6,054,472 and U.S. Patent No. 6,344,465. After the protecting group on the imidazole of 268.15 is removed, the phosphonate containing moiety is introduced to the imidazole to provide compounds of the invention. A specific compound of the invention can be prepared as follows.



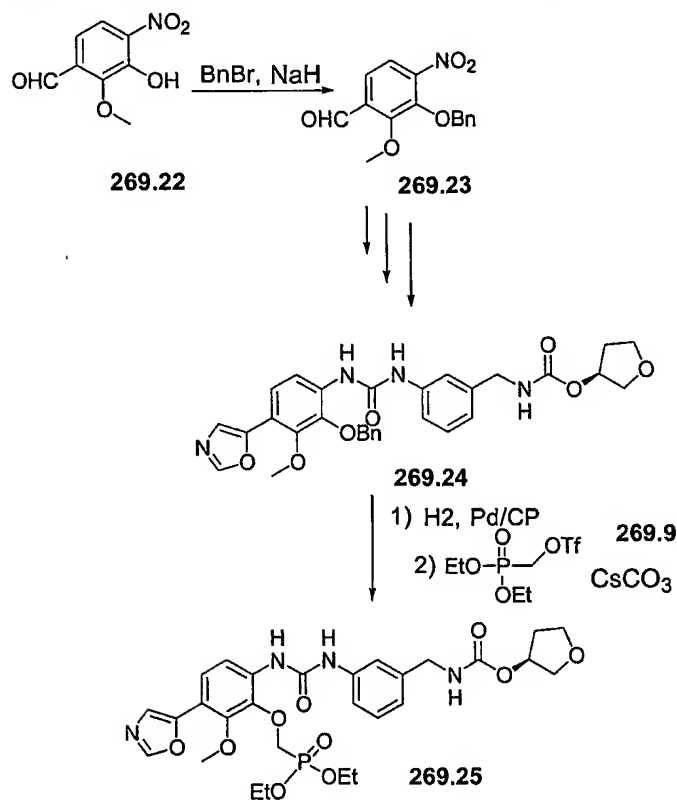
Compound **268.15** is treated with tetrabutylammonium fluoride in THF in reflux condition and the resulting **268.16** is alkylated with **268.9** using sodium hydride as a base to obtain two isomers **268.17** and **268.18**, which are separated by chromatography.

Example 269



Representative compounds of the invention can be prepared as illustrated above. Tetrasubstituted benzene derivatives are obtained by literature

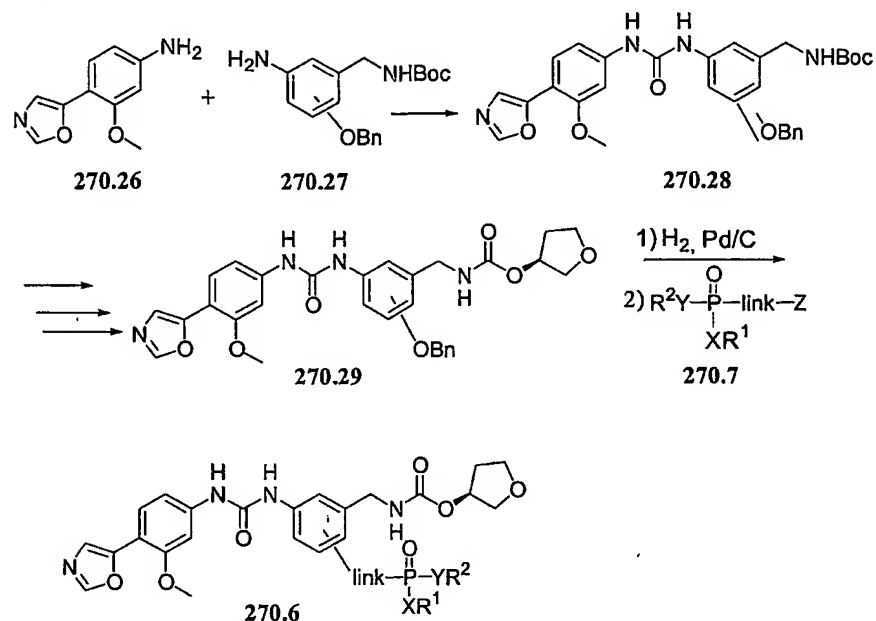
procedures (Ichikawa and Ichibagase *Yakugaku Zasshi* 1963, 83, 103; Norio, A. *et al. Tetrahedron Lett.* 1992, 33(37), 5403). After the phenolic OH is protected with a suitable protecting group, for example benzyl group, the compound **269.21** is synthesized by the same procedure described in U.S. Patent No. 6,054,472, and U.S. Patent No. 6,344,465. After the protecting group is removed, the phosphonate containing moiety is introduced to the phenolic OH using the phosphonate reagent **269.7**, bearing a suitable leaving group. A specific compound of the invention can be prepared as follows.



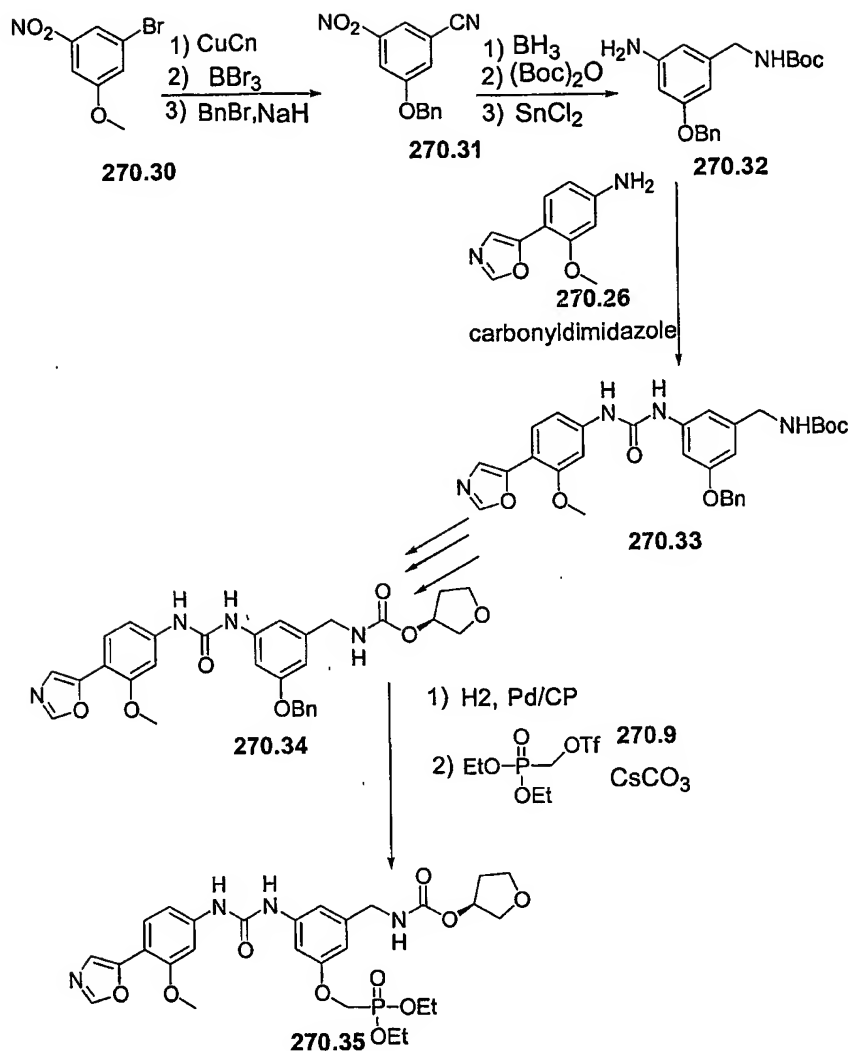
For example, a solution of **269.22**, which is obtained by the procedure of Norio et al. (*Tetrahedron Lett.* 1992, 33(37), 5403), is treated with sodium hydride and one equivalent of benzyl bromide in DMF to get **269.23**. Compound **269.23** is converted to **269.24** by a series of steps such as those reported in U.S. Patent No. 6,054,472, and U.S. Patent No. 6,344,465. After the benzyl protecting group of **269.24** is removed by catalytic hydrogenation, a phosphonate bearing moiety is attached by alkylation of the resulting phenol in

DMF using sodium hydride and one equivalent of (trifluoromethanesulfonyl-oxy)methylphosphonic acid diethyl ester **269.9** to give **269.25**.

Example 270



Representative compounds of the invention can be prepared as illustrated above. Compound **270.26** is treated with carbonyldiimidazole or triphosgene followed by the compound **270.27**, which has a handle to attach phosphonate moiety. Compound **270.27** bearing an extra substituent is synthesized from the tri substituted phenol with a cyano and a nitro groups, which is either commercially available or by literature procedures (Zolfigol, M. A. *et al. Indian J. Chem. Sect. B* **2001**, *40*, 1191; De Jongh, R. O. *et al. Recl. Trav. Chim. Pays-Bas* **1968**, *87*, 1327). The resulting **270.28** is converted to **270.29** using procedures similar to those described in U.S. Patent No. 6,054,472, and U.S. Patent No. 6,344,465. The phosphonate moiety of **270.6** is attached after deprotection of the benzyl group of **270.29**.

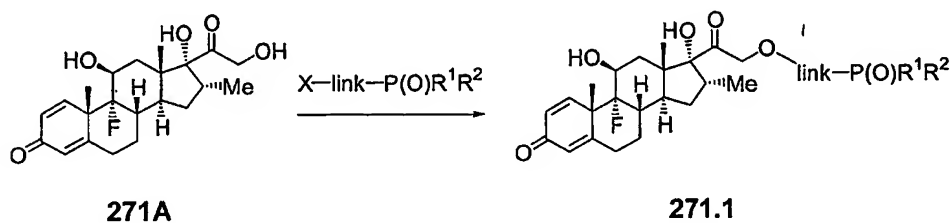


For example, the bromine substituent of compound 270.30 is substituted with cyano group by the procedure of De Jongh, R. O. *et al.* (*Recl. Trav. Chim. Pays-Bas* 1968, 87, 1327) and the methoxy group is converted to benzyloxy group as a protecting group, which affords compound 270.31. After selective reduction of cyano to aminomethyl group by borane, the amino group is protected with Boc group and then the reduction of the nitro group using tin (II) chloride generates compound 270.32. This substituted aniline 270.32 is then treated with a reaction mixture of the compound 270.26 and carbonyldiimidazole, as described in U.S. Patent No. 6,054,472, and U.S. Patent No. 6,344,465, to form the urea 270.33. Compound 270.33 is converted to 270.34. Deprotection of the benzyl group using catalytic hydrogenation

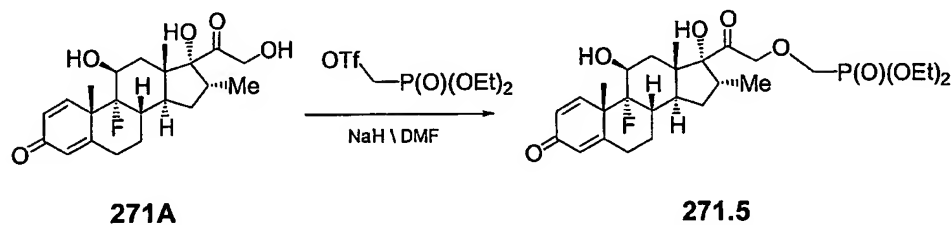
followed by attachment of a phosphonate moiety using **270.9** in the presence of cesium carbonate produces compound **270.35**.

Example 271

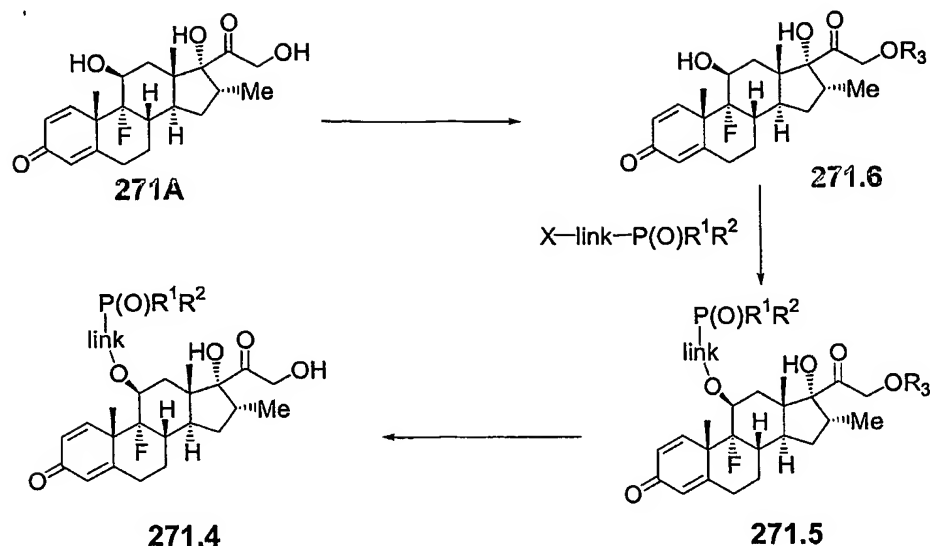
The syntheses of phosphonate compounds of the invention and of intermediate compounds necessary for their synthesis are illustrated herein. Derivatization at the C-21 hydroxy group is accomplished through alkylation of dexamethasone **271A** with the appropriate phosphonate, furnishing analogs shown herein.



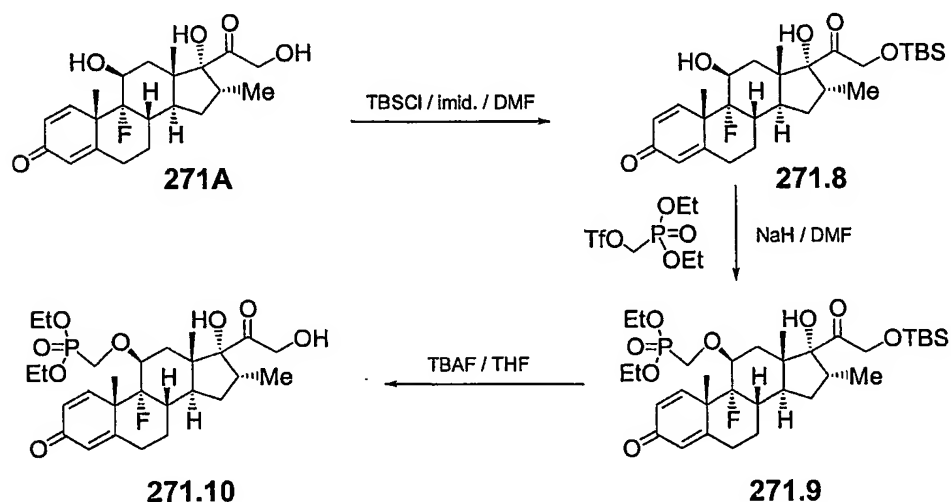
Derivatization at the C-21 hydroxy group is accomplished through alkylation of dexamethasone **271A** with the appropriate phosphonate, furnishing analogs of the type **271.2**.



After sodium hydride extraction of the primary hydroxy proton in **271A**, diethyl phosphonate triflate is added to afford ether **271.5**.

Example 272

Synthesis of C-21 phosphonate analogs having formula, **271.4**, is shown herein. Protection this time of dexamethasone **271A** at the less hindered site furnishes alcohol, **271.5**, which is alkylated at the only exposed hydroxy group with the appropriate phosphonate. Removal of the protecting groups completes the construction of analog, **271.4**.

Example 272

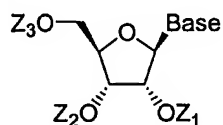
Dexamethasone, **271A**, is protected as its silyl ether using the standard TBSCl and imidazole conditions. (*J. Am. Chem. Soc.*, **1972**, *94*, 6190; however,

harsher conditions should allow for bis-protection. After alkylating with the diethyl phosphonate triflate, the resulting intermediate, **271.9**, is treated with TBAF to give the desired phosphonate, **271.10**.

Example 273

A number of compounds of the general structure **273A** can either be prepared using procedures described in the literature, or be purchased from commercial sources. The following are good sources for information on the art of preparing a variety of compounds of the general structure **273A**, Townsend, *Chemistry of Nucleosides and Nucleotides*, Plenum Press, 1994; and Vorbruggen and Ruh-Pohlenz, *Handbook of Nucleoside Synthesis*, John Wiley & Sons, Inc., 2001.

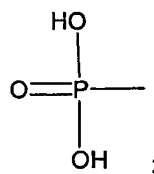
In the examples the compounds of the invention, described herein, the substituents have the following general formula **273A**:



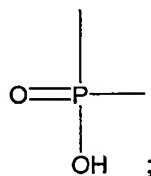
273A

wherein one or more of the Z substituents have been substituted with an A^0 group; and wherein:

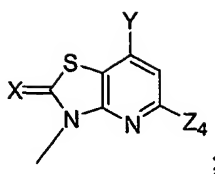
Z_1 and Z_2 are independently selected from hydrogen, or a C_1 - C_{18} acyl, and Z_3 is H, a C_1 - C_{18} acyl, or



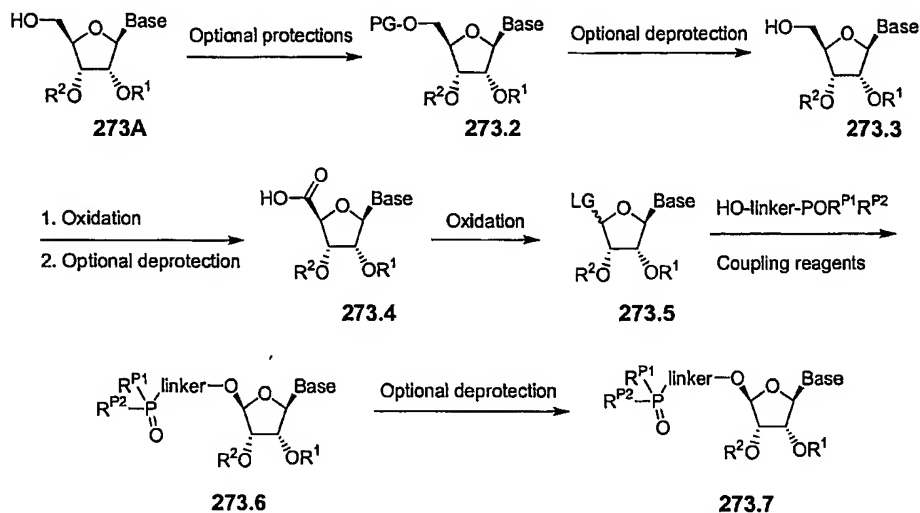
or Z_1 is hydrogen, and together Z_2 and Z_3 are



base is



wherein X and Y are independently O or S; Z^4 is hydrogen, amino, hydroxy, or a halogen selected from Cl and Br.



More specifically, the preparation of generic structure, 273A are described in Nagahara, et al *J. Med. Chem.*; 33; 1990; 407-415. The structure 273.2 is described in Kini, et al *J. Med. Chem.*; 34; 1991; 3006-3010. The two patents cited above also provided examples of the synthesis of compound, 273A.

The core components of this reaction sequence are the transformation of compound from 273.3 to 273.6. Appropriate oxidant(s) can convert the primary alcohol (5'-hydroxy) shown in 273.3 to a carboxylic acid or its corresponding ester. In the case of an ester, an additional deprotection step will give the carboxylic acid, 273.4. A variety of oxidation procedures exist in the literature and can be utilized here. These include but are not limited to the following methods: (i) pyridinium dichromate in Ac_2O , t-BuOH, and dichloromethane producing the t-butyl ester, followed by a deprotection using reagent such as trifluoroacetic acid to convert the ester to the corresponding carboxylic acid (see Classon, et al., *Acta Chem. Scand. Ser. B*; 1985, 39, 501-504. Cristalli, et al., *J. Med. Chem.*, 1988, 31, 1179-1183.); (ii) iodobenzene diacetate and 2,2,6,6-tetramethyl-1-piperidinyloxy, free radical (TEMPO) in acetonitrile, producing

the carboxylic acid (See Epp, *et al*; *J. Org. Chem.* 64; 1999; 293-295. Jung *et al*; *J. Org. Chem.*; 66; 2001; 2624-2635.); (iii) sodium periodate, ruthenium(III) chloride in chloroform producing the carboxylic acid (see Kim, *et al*, *J Med. Chem.* 37; 1994; 4020-4030. Homma, *et al*; *J. Med. Chem.*; 35; 1992; 2881-2890); (iv) chromium trioxide in acetic acid producing the carboxylic acid (see Olsson *et al*; *J. Med. Chem.*; 29; 1986; 1683-1689. Gallo-Rodriguez *et al*; *J. Med. Chem.*; 37; 1994; 636-646); (v) potassium permanganate in aqueous potassium hydroxide producing the carboxylic acid (see Ha, *et al*; *J. Med. Chem.*; 29; 1986; 1683-1689. Franchetti, *et al*; *J. Med. Chem.*; 41; 1998; 1708-1715.) (vi) nucleoside oxidase from *S. maltophilia* to give the carboxylic acid (see Mahmoudian, *et al*; *Tetrahedron*; 54; 1998; 8171-8182.)

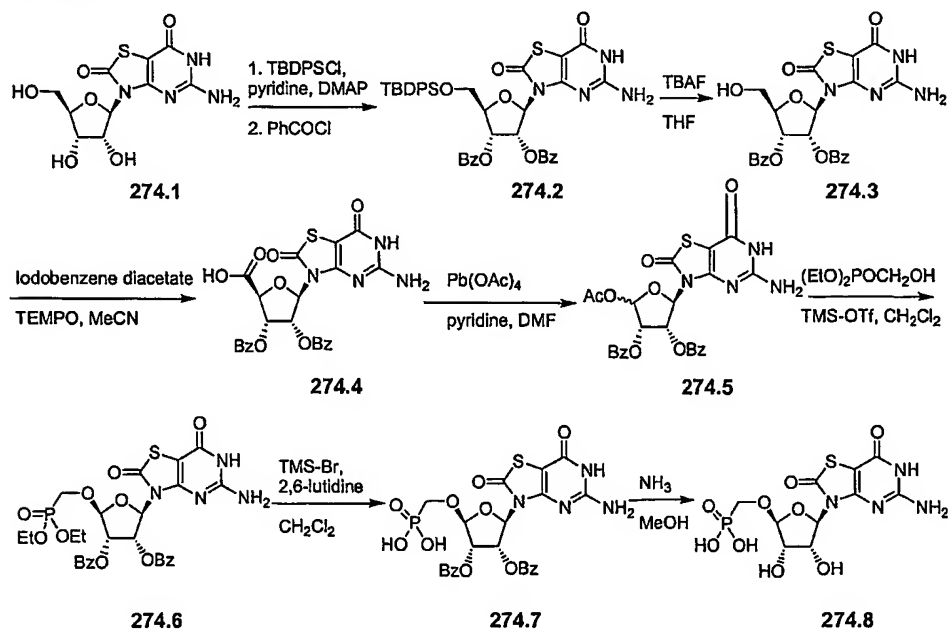
The preparation of compound 273.5 starting with compound 273.4 using lead(IV) tetraacetate (Lv = OAc) was described by Teng *et al*; *J. Org. Chem.*; 59; 1994; 278-280 and Schultz, *et al*; *J. Org. Chem.*; 48; 1983; 3408-3412. When lead(IV) tetraacetate is used together with lithium chloride (see Kochi, *et al*; *J. Am. Chem. Soc.*; 87; 1965; 2052), the corresponding chloride is obtained (273.5, Lv = Cl). Lead(IV) tetraacetate in combination with N-chlorosuccinimide can produce the same product (273.5, Lv = Cl) (see Wang, *et al*; *Tet. Asym.*; 1; 1990; 527 and Wilson *et al*; *Tet. Asym.*; 1; 1990; 525). Alternatively, the acetate leaving group (Lv) can also be converted to other leaving group such as bromide by treatment of trimethylsilyl bromide to give 273.5 ((see Spencer, *et al*; *J. Org. Chem.*; 64; 1999; 3987-3995).

The coupling of 273.5 (Lv = OAc) with a variety of nucleophiles were described by Teng *et al*; *Synlett*; 1996; 346-348 and U.S. Patent No. 6,087,482; Column 54 line 64 to Column 55 line 20. Specifically, the coupling between 273.5 and diethyl hydroxymethylphosphonate in the presence of trimethylsilyl trifluoromethanesulfonate (TMS-OTf) was described. It can be envisioned that other compounds with the general structure of HO-linker-POR^{P1}R^{P2} can also be used so long as the functional groups in these compounds are compatible with the coupling reaction conditions. There are many examples in the published literature describing the coupling of 273.5 (Lv = halogen) with a variety of alcohols. The reactions can be facilitated with a number of reagents, such as silver(I) salts (see Kim *et al*; *J. Org. Chem.*; 56; 1991; 2642-2647, Toikka *et al*;

J. Chem. Soc. Perkins Trans. 1; 13; 1999; 1877-1884), mercury(II) salts (see Veeneman et al; *Recl. Trav. Chim. Pays-Bas*; 106; 1987; 129-131), boron trifluoride diethyl etherate (see Kunz et al; *Hel. Chim. Acta*; 68; 1985; 283-287), Tin(II) chloride (see O'Leary et al; *J. Org. Chem.*; 59; 1994; 6629-6636), alkoxide (see Shortnacy-Fowler et al; *Nucleosides Nucleotides*; 20; 2001; 1583-1598), and iodine (see Kartha et al; *J. Chem. Soc. Perkins Trans. 1*; 2001; 770-772). These methods can be selectively used in conjunction with different methods in forming 273.5 with various leaving groups (Lv) to produce 273.6.

The transformations from 273.1 to 273.2, from 273.2 to 273.3, and from 273.6 to 273.7 are intended to allow the core components of the transformations (from 273.3 to 273.6) to occur while preserving the functional groups already exist in the compound structures. Thus, the syntheses may require the introduction and removal of protecting groups from a compound is a commonly practiced art in organic synthesis. It should be understood that in the transformation 273.6 to 273.7, R^{P1} and R^{P2} do not need to remain unchanged. The final form of R^{P1} and R^{P2} can be selected from a variety of possible structures.

Example 274

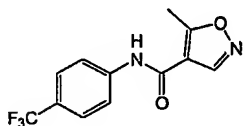


Compound **274.1** is prepared using the method described in the patent application WO 01/90121 (table at page 115). The 5'-hydroxyl in **274.1** is protected as a *tert*-butyldimethylsilyl (TBDMS) ether. The 2'- and 3'- hydroxyl groups can be protected as bezoyl (Bz) esters to give **274.2**. The 5'-hydroxyl can then be deprotected to give **274.3**. Oxidation using iodobenzene diacetate and 2,2,6,6-tetramethyl-1-piperidinyloxy, free radical (TEMPO) convert the primary alcohol to the corresponding acid **274.4**. Further oxidation of **274.4** using lead tetraacetate can produce **274.5**. Coupling between **274.5** and diethyl hydroxy-methyl-phosphonate (available from Sigma-Aldrich, Cat. No. 39,262-6) effected by TMS-OTf can afford **274.6**. Treating **274.6** with TMS-Br converts the phosphodiester to the corresponding phosphonic acid **274.7**. Deprotection of the 2'- and 3'- hydroxyl gives **274.8** as an example of the generic structure A, where Base is an 7-thia-8-oxo-guanosine, R^1 , R^2 , R^{P1} and R^{P2} are hydrogen, linker is a methylene group.

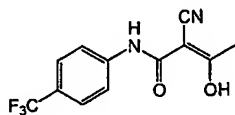
The phosphonic acids in **274.7** and **274.8** are used as examples for illustration purpose. Other forms of phosphonates can be access via the phosphonic acid, or other forms, such as the corresponding diesters as described herein.

Example 275

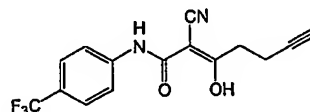
Leflunomide (structure below, together with its active metabolite) (see U.S. Patent No.4,284,786) is a derivative of isoxazole. Representative compounds of the invention can be prepared as illustrated above using procedures similar to those described in *J. Med. Chem.* 1996, 39, 4608. Their structures are shown below.



Leflunomide

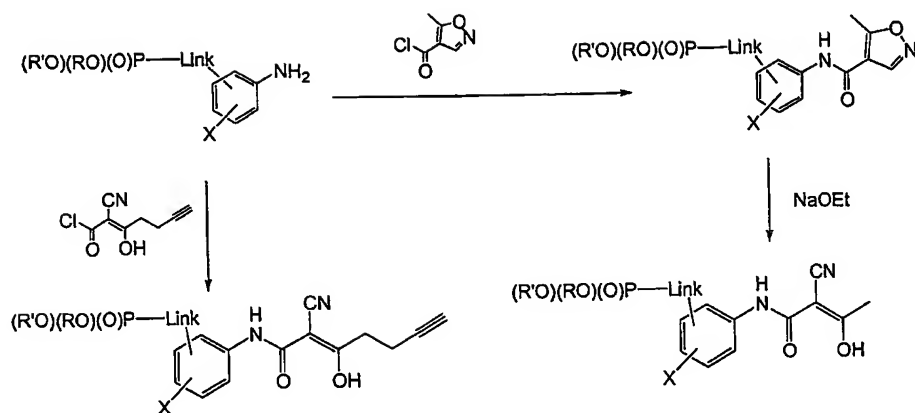


A771726 (active metabolite)

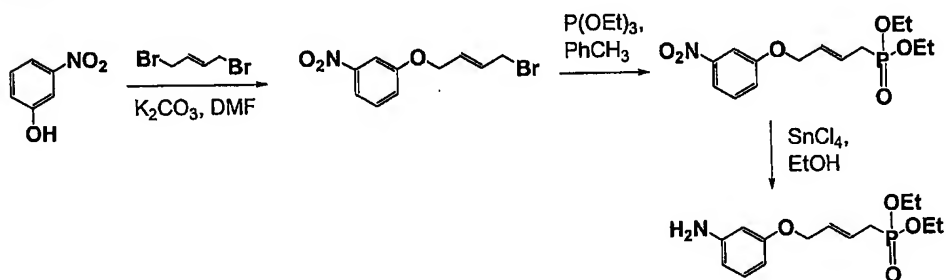
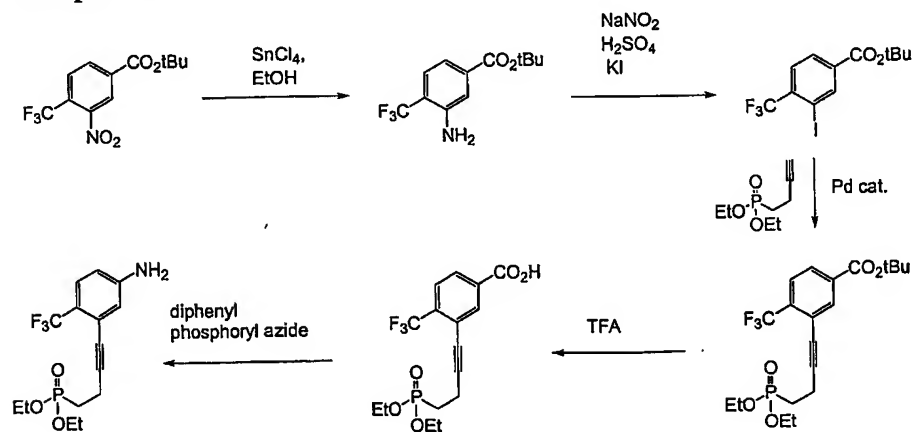


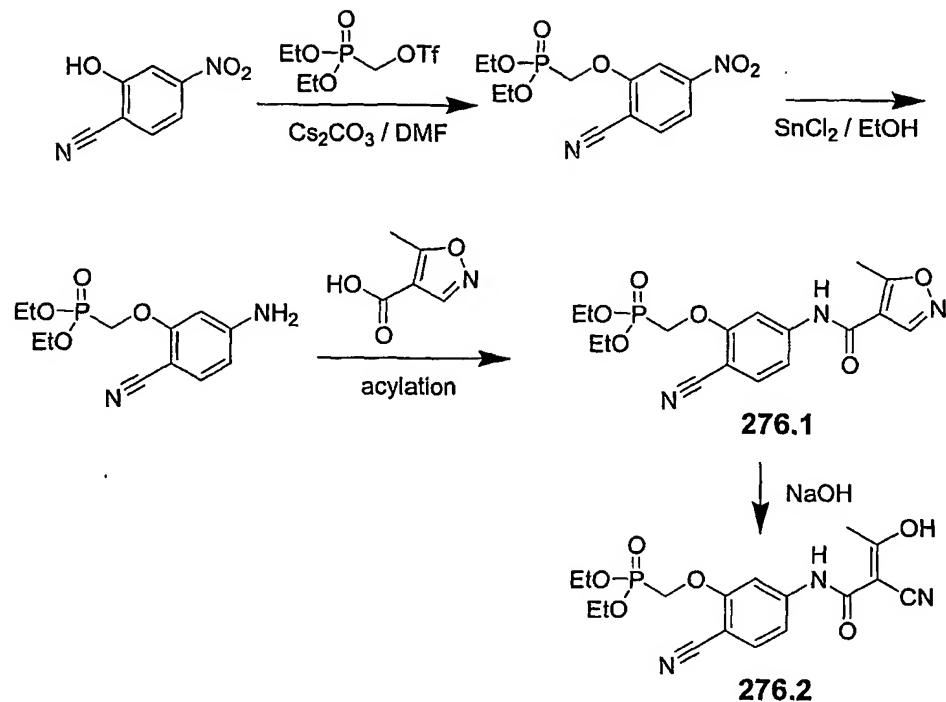
FK778 (MNA-715)

Synthetic methodology towards compounds such as these is described by Westwood et al, *J. Med. Chem.*, 1996, 39, 4608-4621, according to the general routes outlined below.

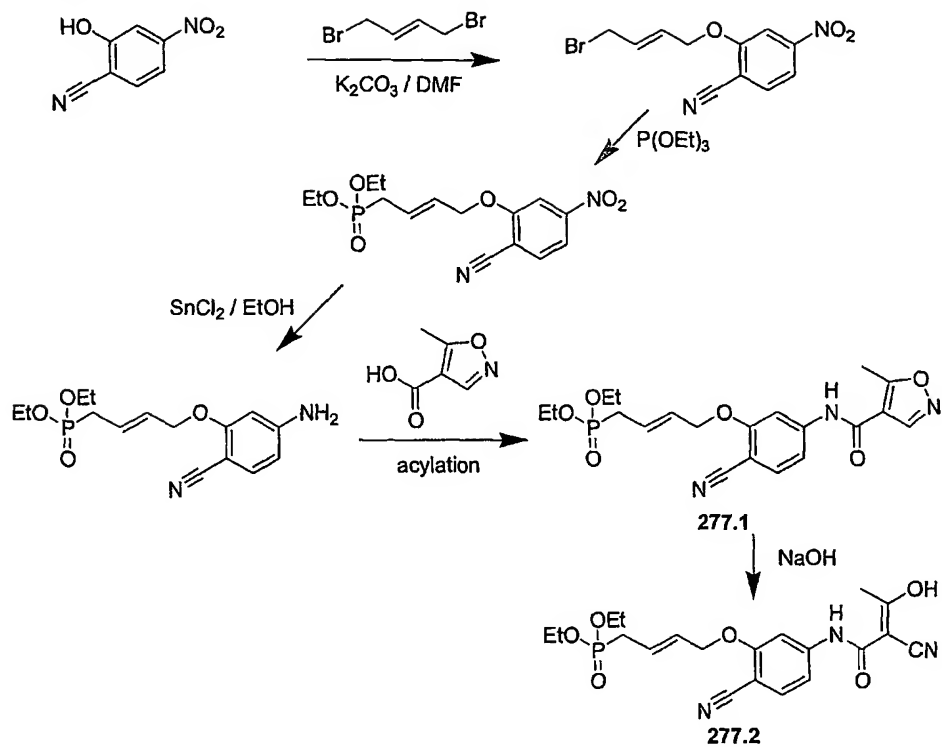
**Example 275A**

The synthesis of suitable phosphonate-containing anilines are shown below.

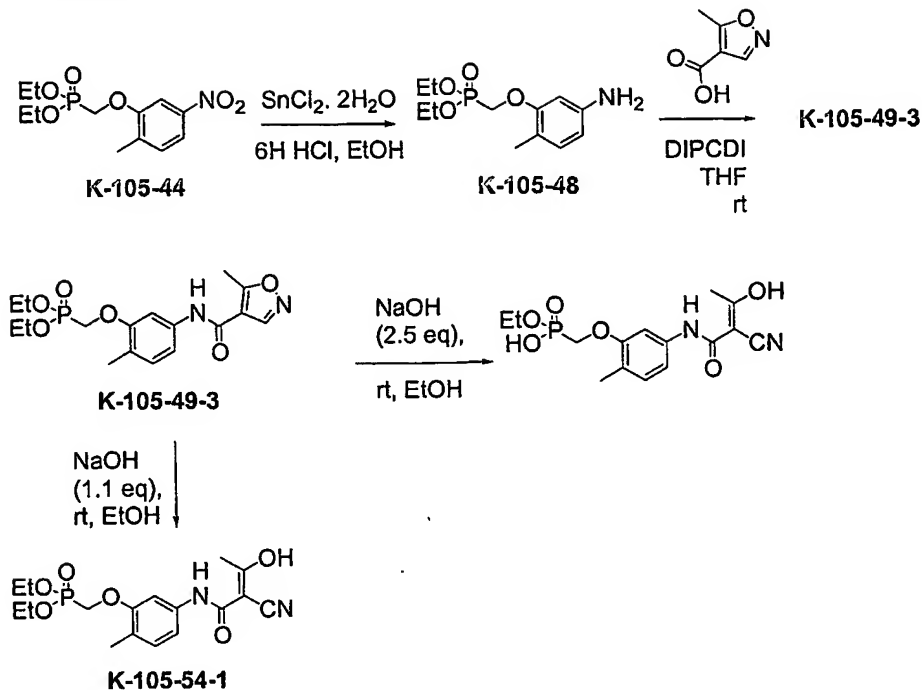
**Example 275B**

Example 276

Representative compounds of the invention can be prepared as illustrated above using procedures similar to those described in *J. Med. Chem.* 1996, 39, 4608. Treatment of compound of the invention **276.1** with base provides compound **276.2** which is also a compound of the invention.

Example 277

Representative compounds of the invention can be prepared as illustrated above. Treatment of compound of the invention 277.1 with base provides compound 277.2 which is also a compound of the invention.

Example 278**Example 278A Synthesis of K-105-44**

2-Methyl-5-nitrophenol (2.00 g, 13.05 mmol) was dissolved in dry DMF (10 mL) under argon atmosphere and cooled to 0°C . Diethylphosphonomethyl-*O*-triflate (4.70 g, 15.66 mmol) and cesium carbonate (6.38 g, 19.58 mmol) were added sequentially. The reaction mixture was stirred at 0°C for 4 hrs. TLC (cyclohexane/EtOAc, 1:1) showed completion of reaction. Deionized water (15 mL) was added and the mixture was extracted with EtOAc (2x50 mL). The organic layer was washed with 1N HCl (20 mL) followed by water (2x20 mL), dried over Na_2SO_4 and concentrated to a semi-solid. Purification by silica gel column chromatography (cyclohexane/EtOAc, 1 : 1) afforded pure compound **K-105-44** as an oil (3.86 g, 97%).

ESI-MS m/z 304 $[\text{M}+\text{H}]^+$.

Example 278B Synthesis of K-105-48

Compound **K-105-44** (2.8 g, 9.24 mmol) was dissolved in 15 mL of absolute ethanol (15 mL) and 6N HCl (2 mL) under an argon atmosphere. Following the addition of $\text{SnCl}_2 \cdot 2\text{H}_2\text{O}$ (5.26 g, 27.72 mmol), the reaction mixture was stirred overnight at room temperature. TLC ($\text{CHCl}_3/\text{MeOH}$, 9 : 1)

showed completion of reaction. The mixture was concentrated to a semi-solid and dissolved in ethyl acetate (30 mL). The ethyl acetate layer was washed with deionized water (10 mL) and satd. NaHCO_3 (10 mL) and dried over Na_2SO_4 . Concentration gave a solid that was used without purification.

ESI-MS m/z 274 $[\text{M}+\text{H}]^+$.

Example 278C Synthesis of K-105-49-3

Crude compound K-105-48 (900 mg, 3.38 mmol) was dissolved in 15 mL of dry THF (15 mL) under an argon atmosphere. Following the addition of 5-methylisoxazole-4-carboxylic acid (381 mg, 3.00 mmol) and diisopropyl carbodiimide (511 μL , 3.30 mmol), the reaction mixture was stirred 6 h at room temperature. TLC ($\text{CHCl}_3/\text{MeOH}$, 9 : 1) showed completion of reaction. The reaction mixture was filtered and the filtrate concentrated to give a solid, which was dissolved in ethyl acetate (25 mL). The solution was washed with deionized water (2x10 mL) and dried over Na_2SO_4 . Concentration gave a solid that was purified by silica gel column chromatography ($\text{CHCl}_3/\text{MeOH}$, 95 : 5) to afford pure compound K-105-49-3 as light yellow solid (680 mg, 55%).

ESI-MS m/z 383 $[\text{M}+\text{H}]^+$.

^1H NMR (300 MHz, CDCl_3): δ 7.11 (1H, s, ArH), 7.06 (2H, s, ArH), 4.29-4.20 (4H, m, OCH_2), 4.14 (2H, d, $J = 10.4$ Hz, OCH_2), 2.76 (3H, s, CH_3), 2.14 (3H, s, CH_3), 1.37 (6H, t, $J = 7.0$ Hz, CH_3).

^{31}P NMR (121.7 MHz, $\text{DMSO}-d_6$ / external H_3PO_4) δ ppm 19.7-20.0 (m)

HPLC: 98 % pure (Sphereclone 5 μL , $\text{H}_2\text{O} : \text{MeCN}$, 20 min linear from 10-90% MeCN , 1.0 mL/min)

Example 278D Synthesis of K-105-54-1

Compound K-105-54-1 (250 mg, 0.65 mmol) was dissolved in 10 mL of absolute ethanol (15 mL) under an argon atmosphere. Following the addition of NaOH (29 mg, 0.72 mmol), the reaction mixture was stirred overnight at room temperature. TLC ($\text{CHCl}_3/\text{MeOH}$, 9 : 1) showed completion of reaction. The reaction mixture was concentrated to a solid and dissolved in ethyl acetate (20 mL). The solution was washed with deionized water (2x10 mL) and dried over Na_2SO_4 . Concentration gave a solid that was purified by silica gel column

chromatography ($\text{CHCl}_3/\text{MeOH}$, 4 : 1), affording pure compound **K-105-54-1** as a solid (188 mg, 75%).

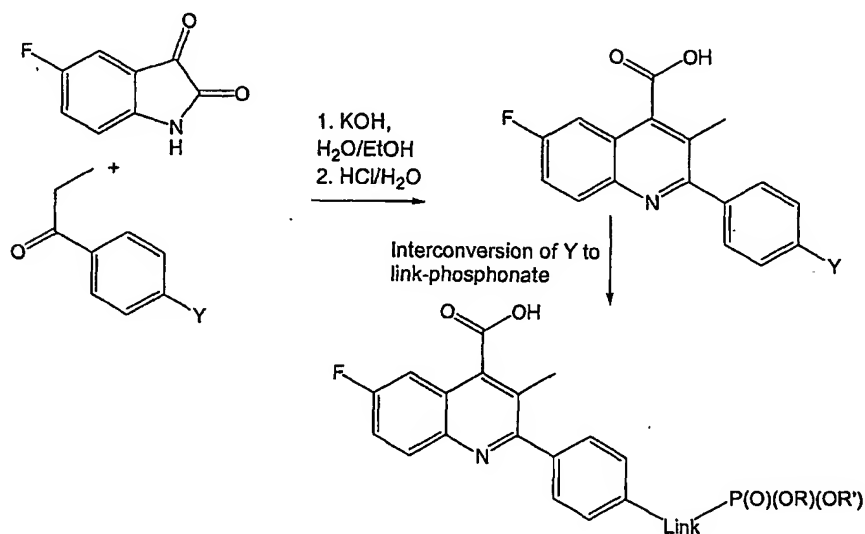
ESI-MS m/z 383 $[\text{M}+\text{H}]^+$.

^1H NMR (300 MHz, $\text{DMSO}-d_6$): δ 7.32 (1H, s, ArH), 6.96 (2H, s, ArH), 4.31 (2H, d, $J = 9.9$ Hz, OCH_2), 4.18-4.08 (4H, m, $2 \times \text{OCH}_2$), 2.08 (3H, s, CH_3), 2.00 (3H, s, CH_3), 1.26 (6H, t, $J = 7.0$ Hz, CH_3).

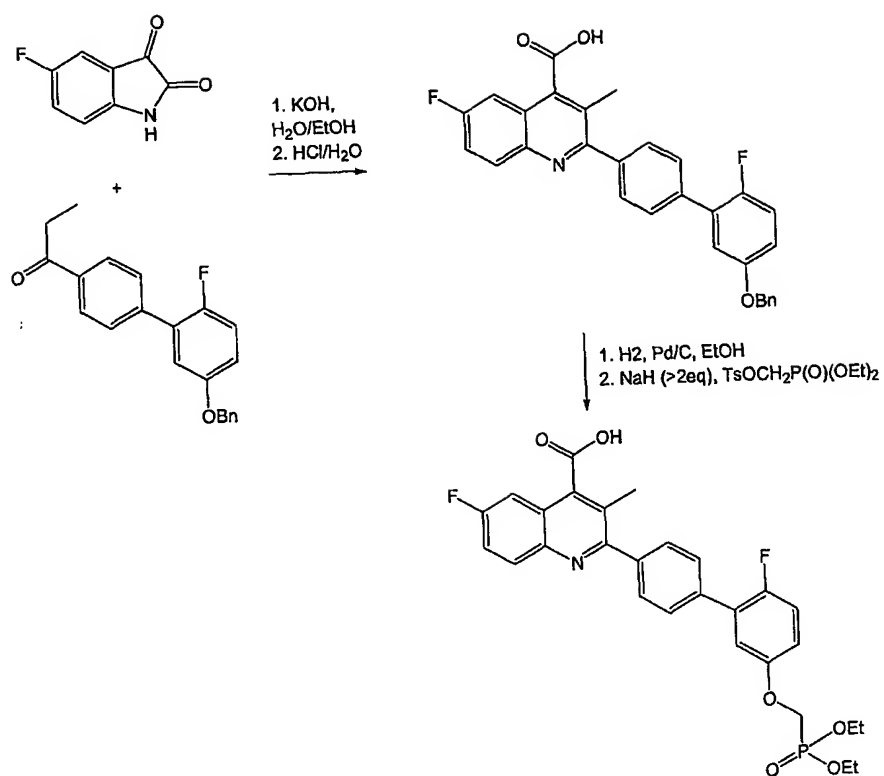
^{31}P NMR (121.7 MHz, $\text{DMSO}-d_6$ / external H_3PO_4) δ ppm 20.0-20.4 (m)

HPLC: 93% pure (Sphereclone 5 μL , H_2O : MeCN , 20 min linear from 10-90% MeCN , 1.0 mL/min)

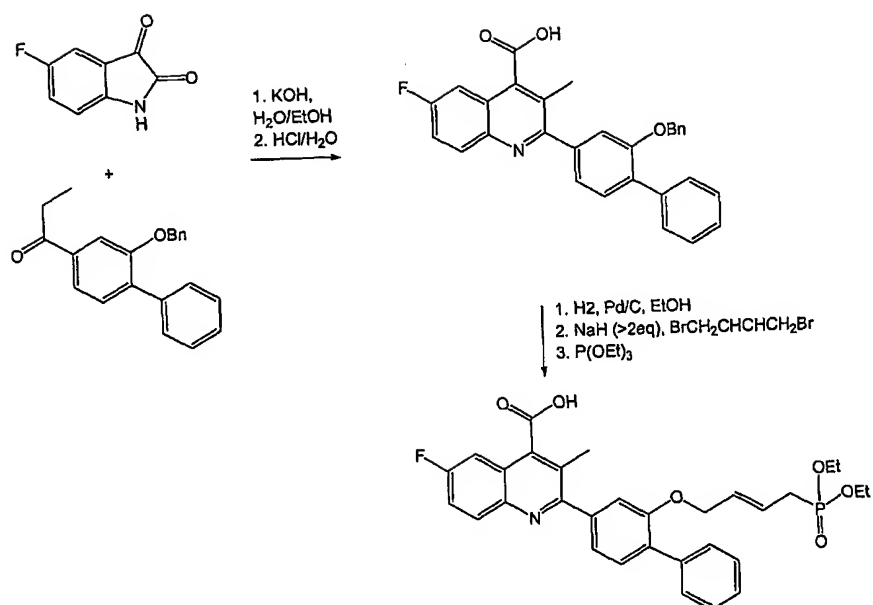
Example 279

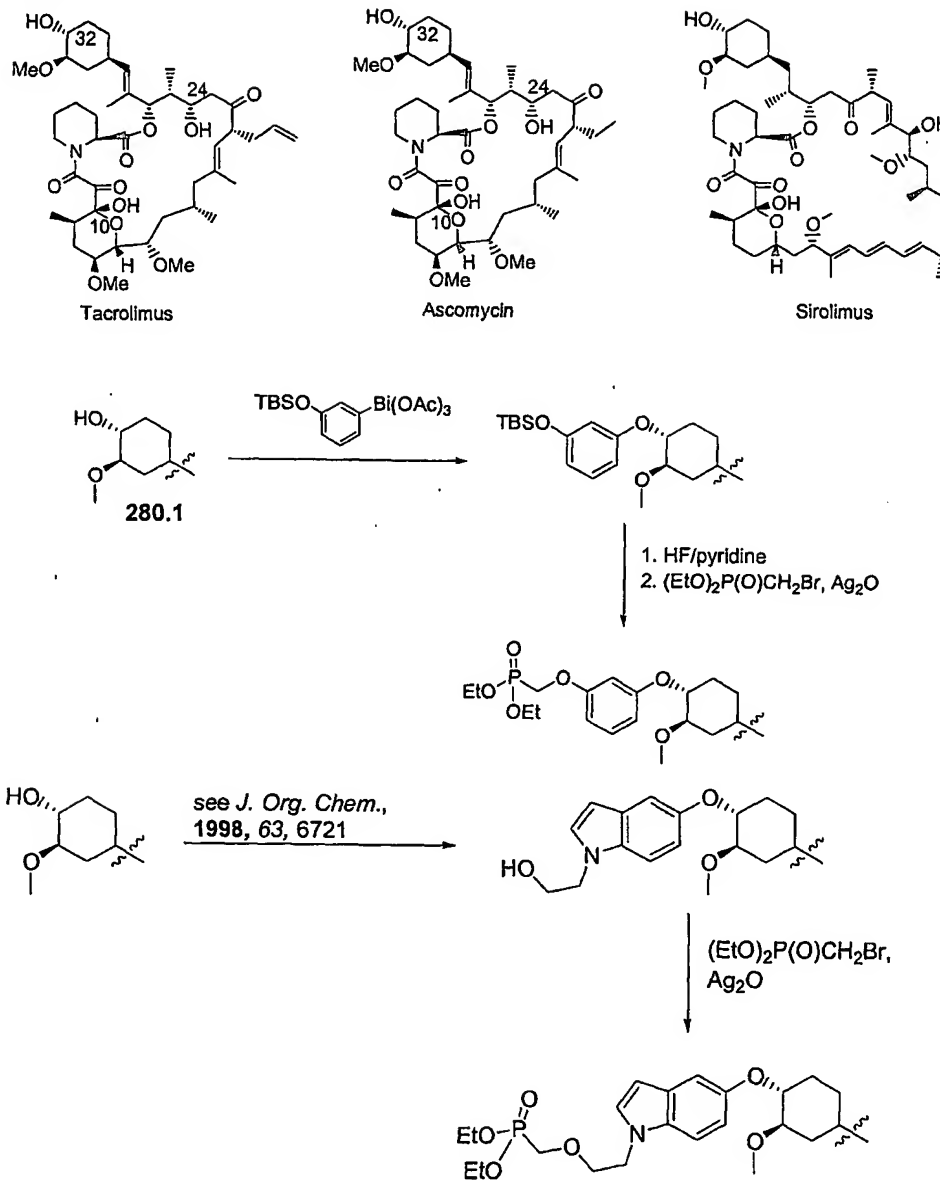


Representative compounds of the invention can be prepared as illustrated above. Specific compounds of the invention can be prepared as illustrated below.

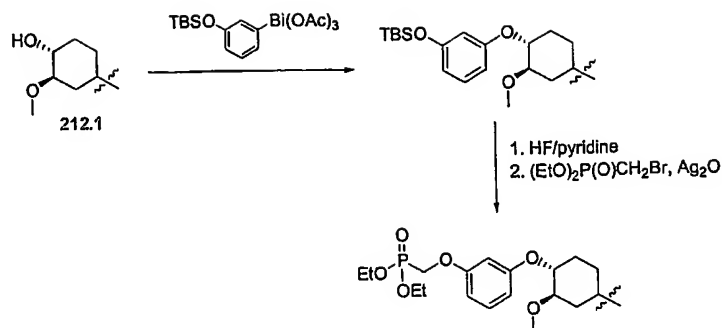


and

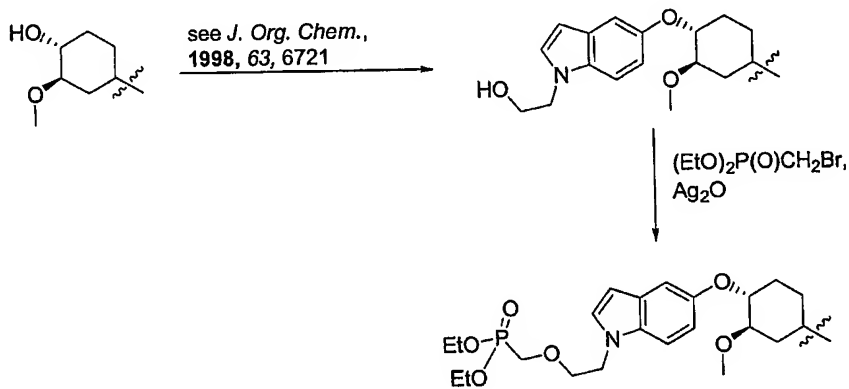
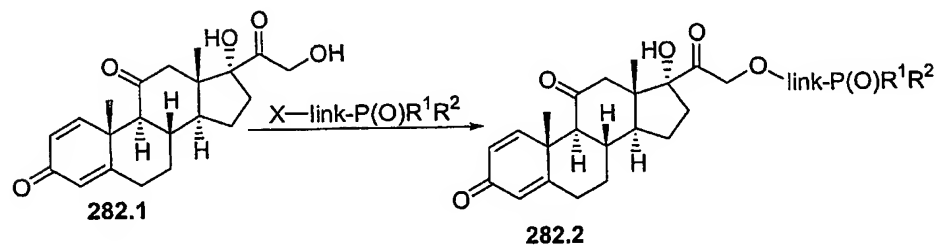


Example 280

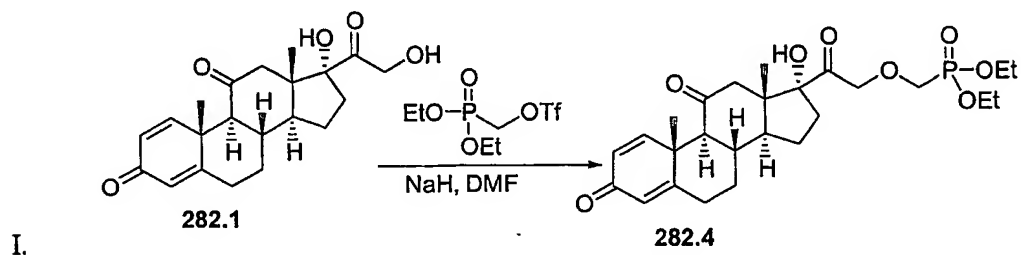
Representative macrolide compounds of the invention, wherein the structure **280.1** is understood to be the compound tacrolimus, ascomycin or sirolimus, can be prepared as illustrated above, for example, using an aryl bismuth reagent such as that shown is described in *Bioorg. Med. Chem. Lett.*, 1995, 5, 1035. Additionally, silver salts have been used to mediate alkylations on immunosuppressive macrolides such as these: see *J. Med. Chem.*, 1998, 41, 1764. Specific compounds of the invention can be prepared as illustrated below.

Example 281

Representative macrolide compounds of the invention, wherein the structure 212.1 is understood to be the compound tacrolimus, ascomycin or sirolimus, can be prepared as illustrated above, for example, using an aryl bismuth reagent such as that shown is described in *Bioorg. Med. Chem. Lett.*, 1995, 5, 1035. Additionally, silver salts have been used to mediate alkylations on immunosuppressive macrolides such as these: see *J. Med. Chem.*, 1998, 41, 1764. Specific compounds of the invention can be prepared as illustrated below.

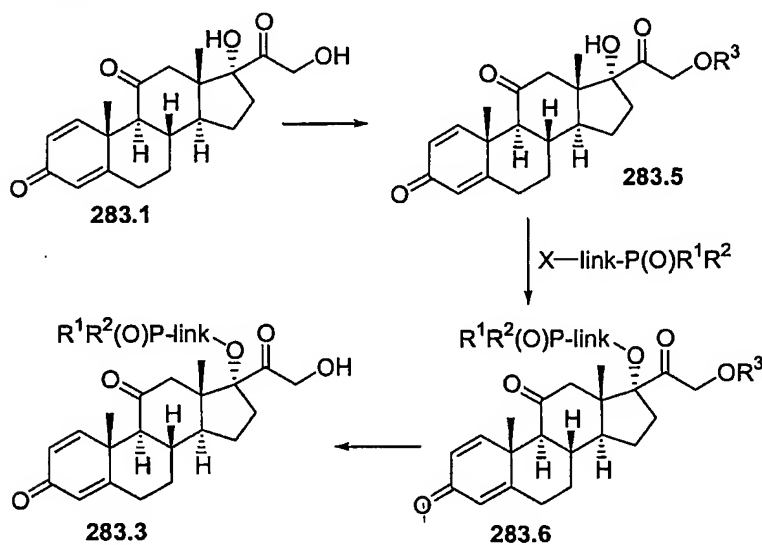
**Example 282**

Representative compounds of the invention can be prepared as illustrated above. Derivatization at the C-21 hydroxy group is accomplished through alkylation of prednisone **282.1** with the appropriate phosphonate to provide compounds of the invention **282.2**. A specific compound of the invention can be prepared as follows.



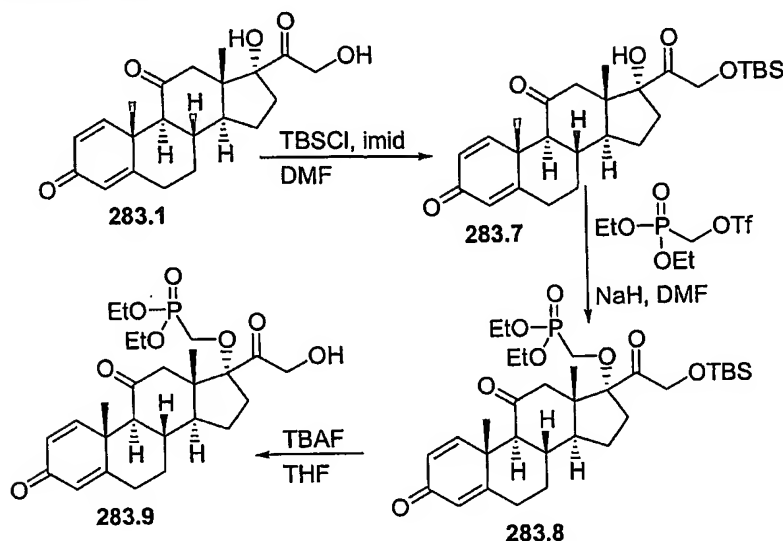
After sodium hydride extraction of the primary hydroxy proton in **282.1**, diethyl phosphonate triflate is added to afford ether **282.4**.

Example 283



Representative compounds of the invention **283.3** can be prepared as illustrated above. Protection of prednisone **283.1** at the less hindered primary site furnishes alcohol **283.5**, which is alkylated at the exposed hydroxy group with the appropriate phosphonate to provide **283.6**. Removal of the protecting

group completes the construction of analog 283.3. A specific compound can be prepared as follows.

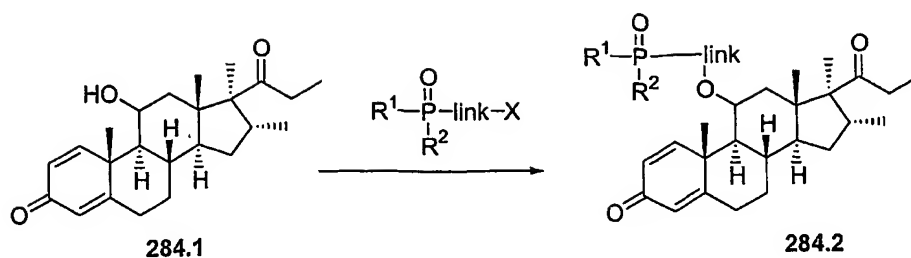


Prednisone 283.1 is mono-protected as its TBS ether 283.7. After alkylating with the diethyl phosphonate triflate, the resulting intermediate 283.8 is treated with TBAF to give the desired phosphonate 283.9.

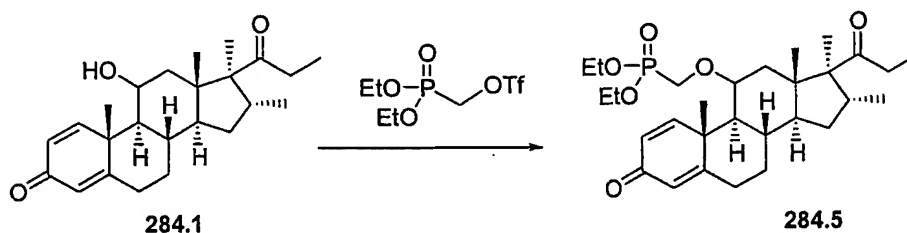
The syntheses of phosphonate compounds of the invention and of intermediate compounds necessary for their synthesis are illustrated herein. Derivatization at the C-21 hydroxy group is accomplished through alkylation of dexamethasone 1 with the appropriate phosphonate, furnishing analogs shown herein.

Example 284

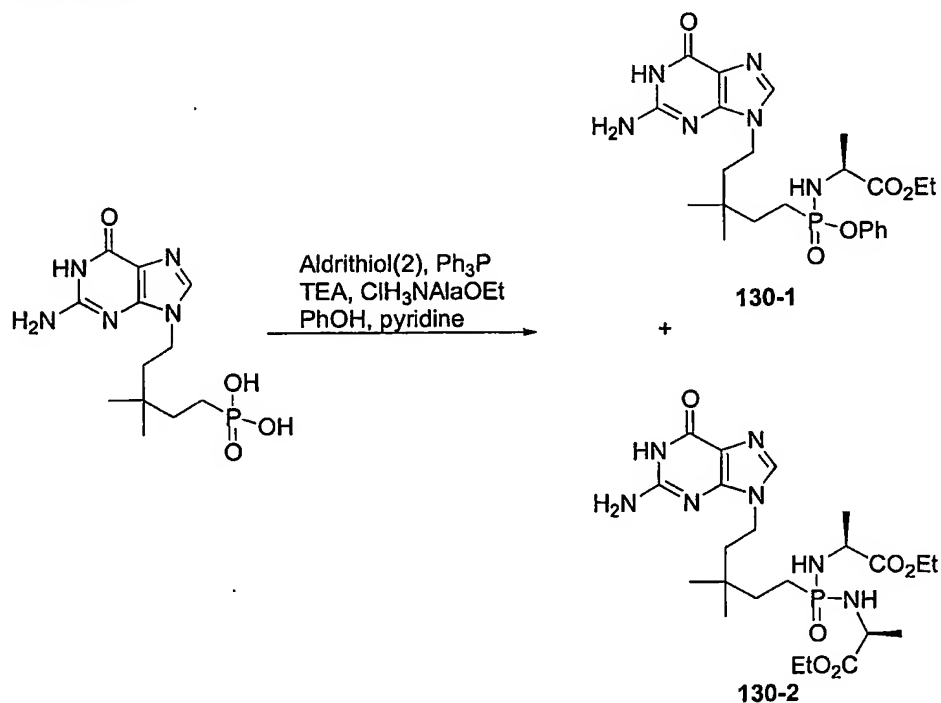
Representative compounds of the invention can be prepared as illustrated above. Derivatization at the C-11 hydroxy group is accomplished through alkylation of rimexolone 284.1 with the appropriate phosphonate, furnishing analogs of formula 284.2. A specific compound of the invention can be prepared as illustrated below.



After sodium hydride extraction of the hydroxy proton in 284.1, diethyl phosphonate triflate is added to afford ether 284.5.



Example 285

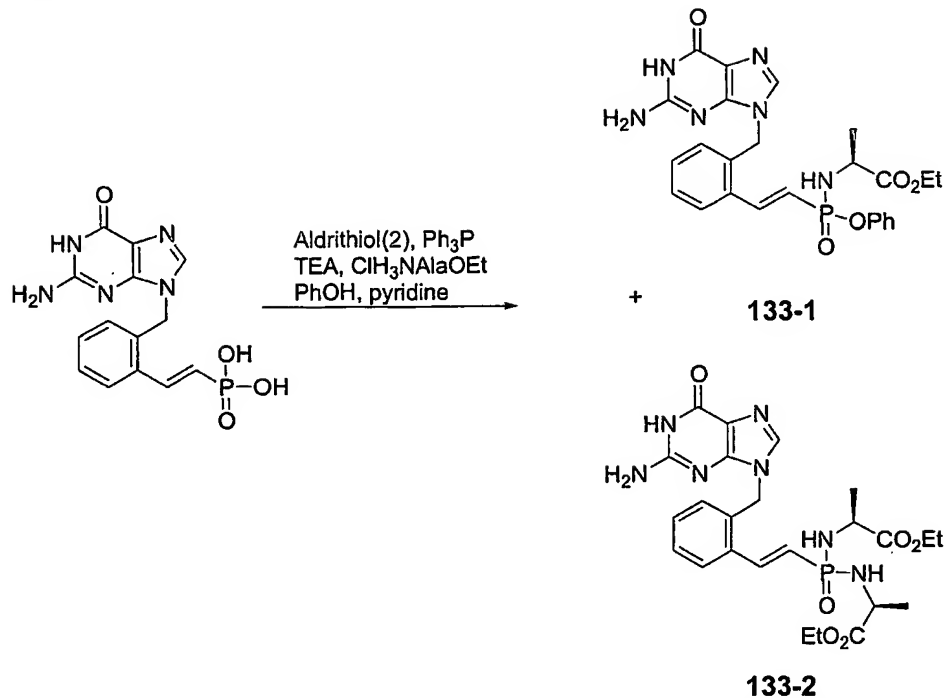


The diacid (100 mg, 0.304 mmol), amino acid (100 mg, 0.651 mmol), phenol (145 mg, 1.54 mmol), and triethylamine (510 μL , 3.66 mmol) were dissolved in pyridine (5 mL). The mixture was heated to 60°C for 5 minutes.

To this reaction mixture was added a solution of triphenylphosphine (560 mg, 2.14 mmol) and Aldrithiol(2) (470 mg, 2.13 mmol) dissolved in pyridine (5 mL). The reaction was then heated at 60°C for 12 hours. The reaction mixture was diluted in EtOAc, washed with H₂O, sat'd NaHCO₃(aq), and brine. The organic layer was dried (MgSO₄), concentrated and purified by chromatography on silica gel (1% MeOH/CH₂Cl₂ → 10% MeOH/CH₂Cl₂) to give monoamidate 130-1 (5 mg, 3%) and bisamidate 130-2 (5 mg, 3%).

For 130-1: ¹H NMR (300 MHz, CD₃OD) δ 7.78 (1H, m), 7.35 (2H, m), 7.20 (3H, m), 4.18-3.95 (5H, m), 2.24-1.90 (2H, m), 1.87-1.62 (4H, m), 1.38-1.18 (6H, m), 1.02 (6H, m); ³¹P NMR (121 MHz, CD₃OD) δ 36.3, 35.3; LC-MS (method: 0.5 min 95% H₂O/5% MeCN → 5 min 0% H₂O/100% MeCN, rt = 2.18 min. MS calc'd for C₂₃H₃₄N₆O₅P (MH⁺): 505.2. Found 505.2.

For 130-2: ¹H NMR (300 MHz, CD₃OD) δ 7.77 (1H, s), 4.23-3.92 (8H, m), 2.04-1.50 (6H, m), 1.42 (3H, d), 1.40 (3H, d), 1.28 (3H, t), 1.22 (3H, t), 1.02 (3H, s), 1.01 (3H, s); ³¹P NMR (121 MHz, CD₃OD) δ 33.9; LC-MS (method: 0.5 min 95% H₂O/5% MeCN → 5 min 0% H₂O/100% MeCN, rt = 1.79 min. MS calc'd for C₂₂H₃₉N₇O₆P (MH⁺): 528.3. Found 528.3.

Example 286

The diacid (25 mg, 0.072 mmol), amino acid (25 mg, 0.16 mmol), phenol (38 mg, 0.40 mmol), and triethylamine (127 μL , 0.911 mmol) were dissolved in pyridine (1.25 mL). The mixture was heated to 60°C for 5 minutes. To this reaction mixture was added a solution of triphenylphosphine (140 mg, 0.534 mmol) and Aldrithiol(2) (119 mg, 0.540 mmol) dissolved in pyridine (1.25 mL). The reaction was then heated at 60°C for 12 hours. Another batch of diacid (12 mg, 0.035 mmol) was treated as described above. The reaction mixtures from both batches were combined and diluted in EtOAc, washed with H_2O , sat'd $\text{NaHCO}_3(\text{aq})$ and brine. The organic layer was dried (MgSO_4), concentrated and purified by chromatography on silica gel (1% MeOH/ $\text{CH}_2\text{Cl}_2 \rightarrow 10\%$ MeOH/ CH_2Cl_2). to give monoamidate **133-1** (3 mg, 8%) and bisamidate **133-2** (8 mg, 20%).

For **133-1**: ^1H NMR (300 MHz, CD_3OD) δ 8.38-8.08 (1H, m), 7.78-7.60 (2H, m), 7.50-7.18 (8H, m), 6.67-6.05 (1H, m), 5.60-5.30 (2H, m), 4.63 (1H, bs), 4.25-3.95 (3H, m), 1.37 (3H, m), 1.18 (3H, m); ^{31}P NMR (121 MHz, CD_3OD) δ 21.5, 20.2; LC-MS (method: 0.5 min 95% H_2O /5% MeCN \rightarrow 5 min 0% H_2O /100% MeCN, rt = 1.98 min. MS calc'd for $\text{C}_{25}\text{H}_{28}\text{N}_6\text{O}_5\text{P}$ (MH^+): 523.2. Found 523.2.

For 133-2: ^1H NMR (300 MHz, CD_3OD) δ 8.15 (1H, dd), 7.72 (1H, s), 7.67 (1H, m), 7.39 (2H, m), 7.28 (1H, m), 6.44 (1H, dd), 5.40 (2H, s), 4.23-3.90 (6H, m), 1.42 (6H, m), 1.27 (3H, t), 1.18 (3H, t); ^{31}P NMR (121 MHz, CD_3OD) δ 19.7; LC-MS (method: 0.5 min 95% H_2O /5% MeCN \rightarrow 5 min 0% H_2O /100% MeCN, rt = 1.86 min. MS calc'd for $\text{C}_{24}\text{H}_{33}\text{N}_7\text{O}_6\text{P}$ (MH^+): 546.2. Found 546.2.

Example 287 Pro-drug cleavage assays

Isolation of PBMC Extracts:

Fresh human PBMCs were obtained from patients undergoing leukapheresis; cells were shipped in plasma and processed within 26 h of draw. Purification was achieved using the Ficoll-Paque method: PBMC cells were harvested by centrifugation at 1200 X g for 5 minutes and washed three times by re-suspension in RBC lysis buffer (155 mM NH_4Cl , 0.1 mM EDTA, 10mM KHCO_3). Washed cells were suspended in lysis buffer (0.2×10^9 cells in 1 ml of 10 mM Tris, pH 7.4, 150 mM NaCl, 20 mM CaCl_2 , 1 mM DTT and 1% NP40) and incubated on ice for 20 minutes. The PBMC crude extract was centrifuged at 1000 X g for 30 min to remove unlysed cells and the supernatant at 100,000 X g for 1h. The 100,000 X g supernatant (PBMC Extract: P0) was harvested, snap frozen in liquid nitrogen and stored at -70°C .

Protocol for measurement of cleavage of prodrugs by PBMC Extracts:

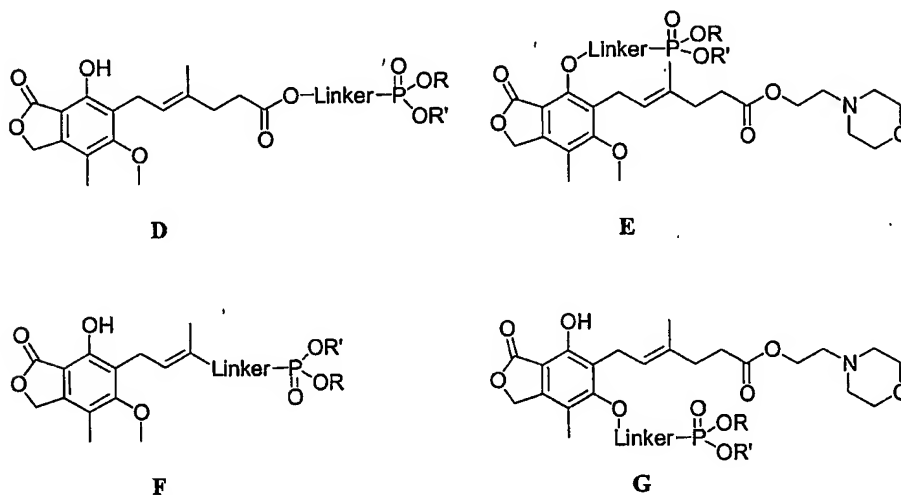
Reaction mixtures contained 25mM MesNa (pH 6.5), 100 mM NaCl, 1 mM DTT, 0.1% NP-40, 30 μM substrate, and varying amounts of enzyme in a final volume of 100 μl . The enzymatic reaction is performed at 37°C for 10–120 minutes and stopped at 3-4 individual time points by adding 180 μl of ice cold methanol. Samples are incubated @ -20°C for 30 min, and centrifuged 13,000 RPM for 30 min (@ 4°C). The supernatant is transferred to a 96 well plate and evaporated under vacuum using a speedvac. The precipitate is dissolved in 100 μl of 20 mM $\text{CH}_3\text{COONH}_4$ + 5% AcCN. The disappearance of pro-drug is measured by HPLC, monitoring at 260nm. The specific activity of the PBMC Extract against the prodrugs tested is defined as: v (cleavage rate) / μg protein = pmoles / min / μg .

Results

Compound	Human PBMC extract specific activity (pmol/min/ μ g)
130-1	3.48
130-2	0.65
133-1	4.9
133-2	0.38

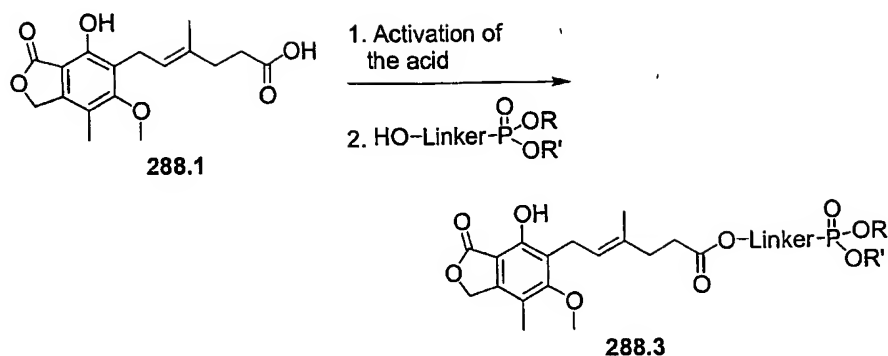
Example 288

Representative compounds of the invention having the following formulae can be prepared as described herein.

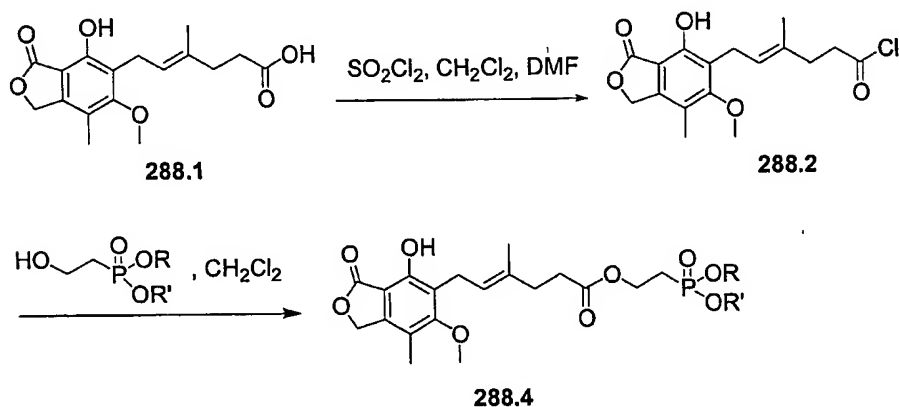


For example, three regions of mycophenolate mofetil can be utilized for the attachment of the phosphonate prodrug as demonstrated by compounds **D**, **E**, and **G** shown above. Also, the carboxylic acid can be replaced with a phosphonic acid as in compound **F**.

Example 288A



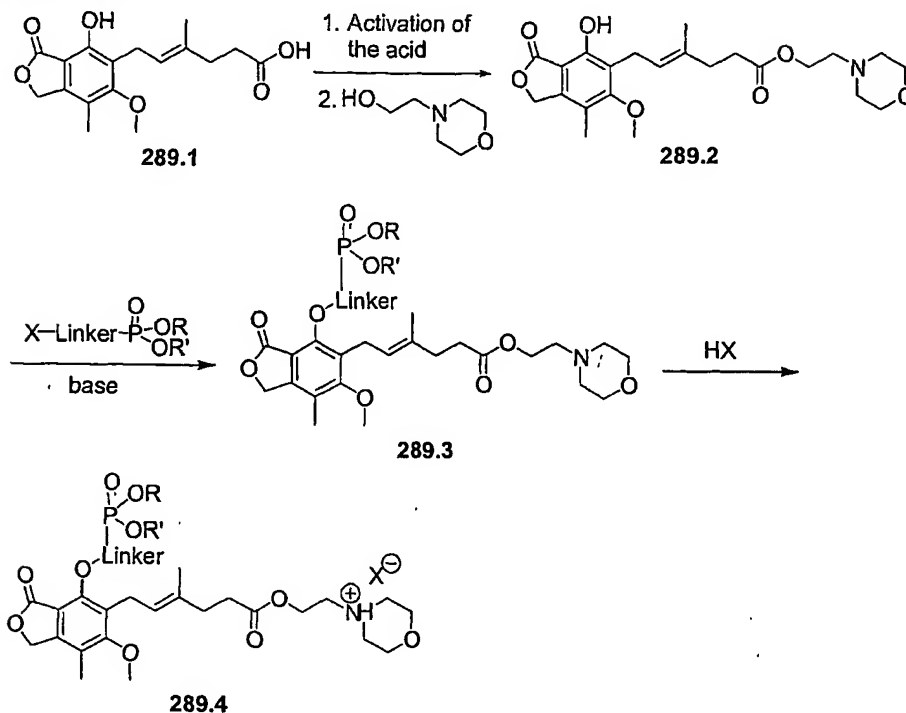
Representative compounds of the invention can be prepared as illustrated above. The morpholino ethyl moiety can serve as a prodrug functionality to improve bioavailability and can be replaced with the phosphonate prodrug handle as shown above. Mycophenolic acid is commercially available, *e.g.*, from Sigma Chemical Company, St. Louis, Mo. Activation of the carboxylic acid **288.1** in the presence of the free phenol, followed by addition of an alcohol carrying the phosphonate group, results in the formation of the desired product **288.3** (U.S. Patent No. 4,786,637). A specific compound of the invention can be prepared as follows.



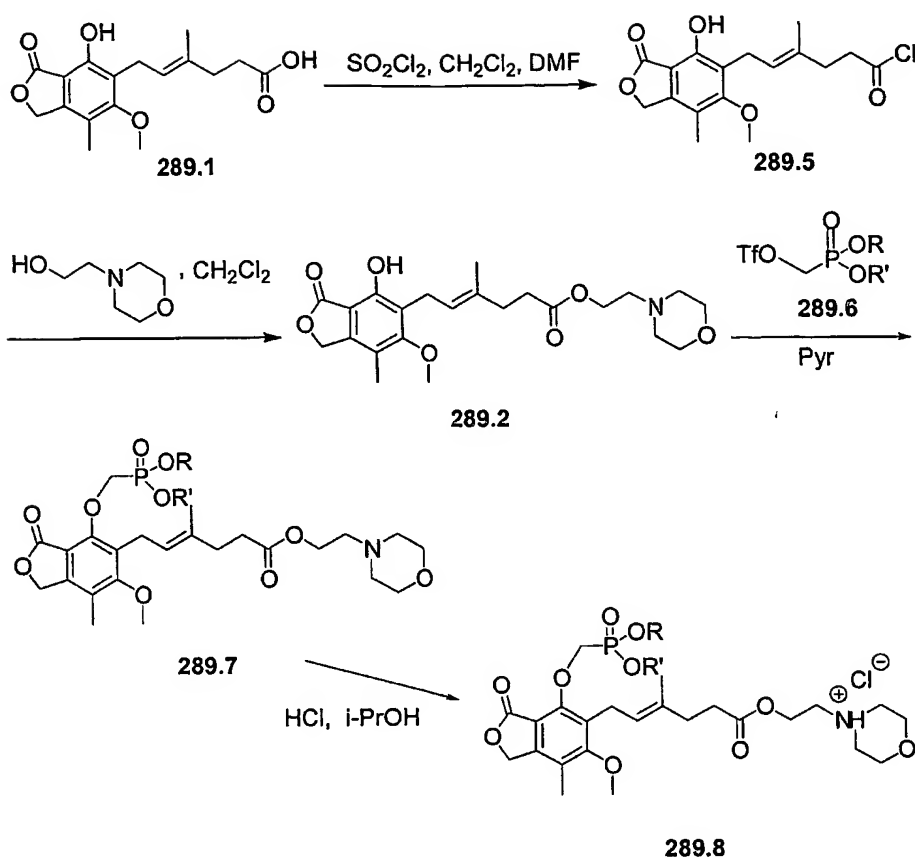
Mycophenolic acid **288.1** is dissolved in dichloromethane. Thionyl chloride is added followed by a catalytic amount of DMF. The reaction mixture is stirred at room temperature for 3 hours, after which the volatile components are removed under vacuum. The phosphonate-alcohol is dissolved in dichloromethane and chilled to about 4 °C on an ice bath. The mycophenolic acid chloride **288.2** is dissolved in dichloromethane and added to the chilled solution. After stirring for 90 minutes at about 4 °C, the reaction mixture is

washed with water and then with aqueous sodium bicarbonate. The organic solution is dried and evaporated to yield the phosphonate **288.4**.

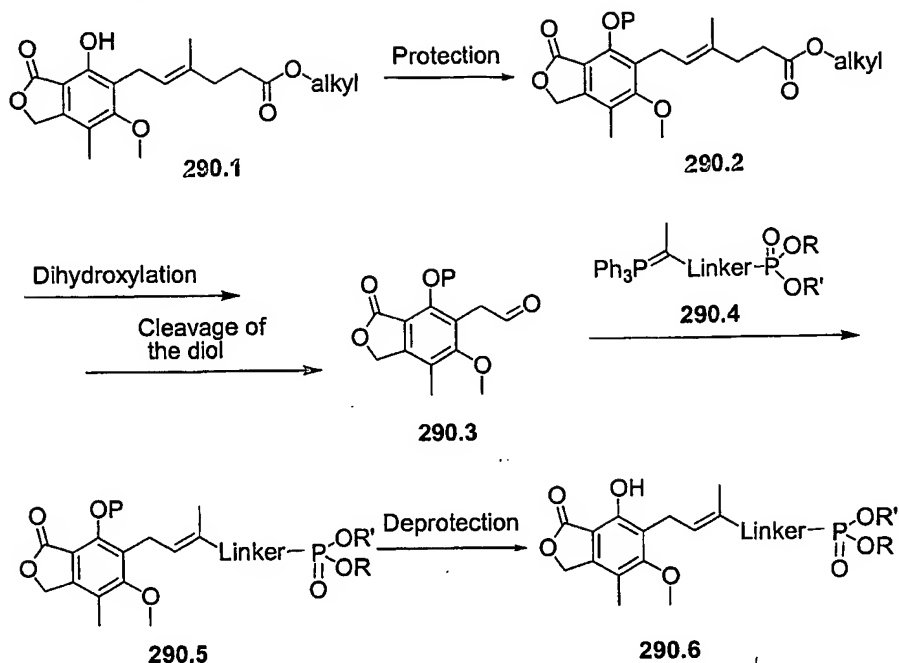
Example 289



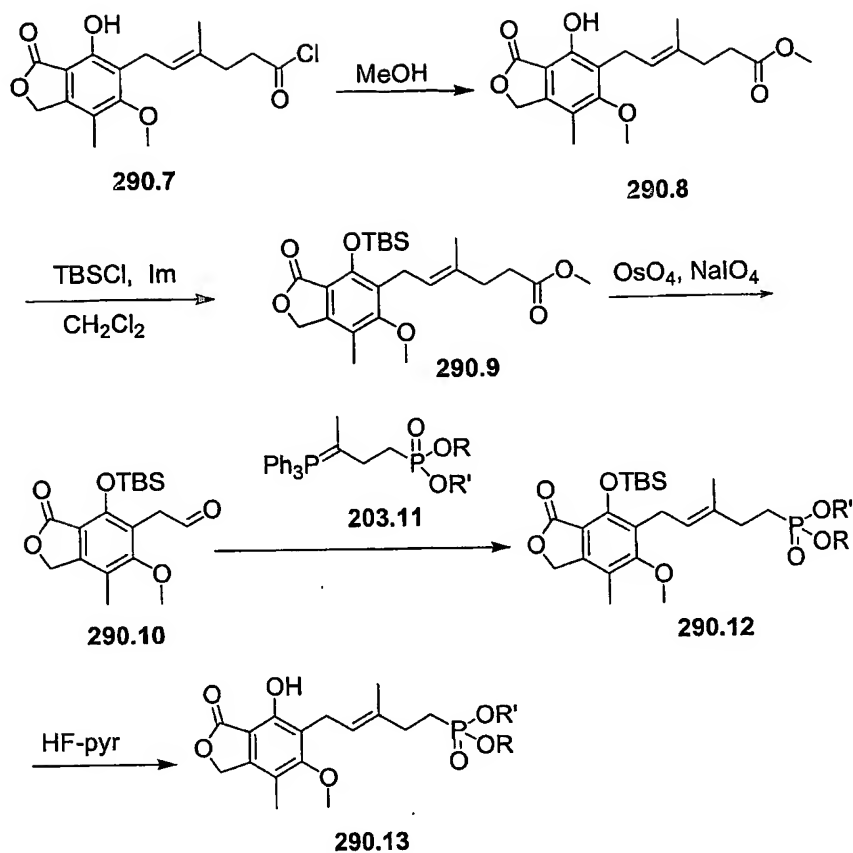
Representative compounds of the invention can be prepared as illustrated above. The C-4 phenol position provides a reactive handle for further analogs as illustrated above. Once the carboxylic acid of **289.1** is blocked by morpholino ethyl, such as in compound **289.2** the phenol can be alkylated under basic conditions. Bases such as pyridine, potassium carbonate, or triethylamine are utilized. Leaving groups such as trifluoromethylsulfonate, mesylate, bromide, or iodide are attached to the phosphonate prodrug subunit and reacted, in the presence of base, with compound **289.2**. Compound **289.3** can either be used directly, or in the form of a salt, compound **289.4**. Among the large number of salts that can be prepared, chloride and bisulfate salts are one particular embodiment of the invention. A specific compound of the invention can be prepared as follows.



Compound 289.5 is prepared similar to compound 289.2 (described in Example 288). A solution of morpholino ethanol in dichloromethane is cooled to about 4 °C. The mycophenolic acid chloride 289.5 is dissolved in dichloromethane and added to the cooled solution. Stirring this solution for about 90 minutes gives compound 289.2. The reaction mixture is washed with water and dried with sodium sulfate. Removal of the solvent provides isolated compound 289.2. Alkylation at the phenolic position of 289.2 is achieved by suspending the compound in pyridine. Triplate 289.6 is added to the solution and the mixture is stirred at room temperature for about 90 minutes. The reaction mixture is poured into water and the product is extracted with ethyl acetate. Removal of the organic layer provides compound 289.7. Hydrochloride salt of 289.7 can optionally be prepared. Compound 289.7 is dissolved in isopropanol and the solution is added to a mixture of hydrogen chloride in isopropanol. The hydrochloride salt 289.8 is collected by filtration and dried under vacuum.

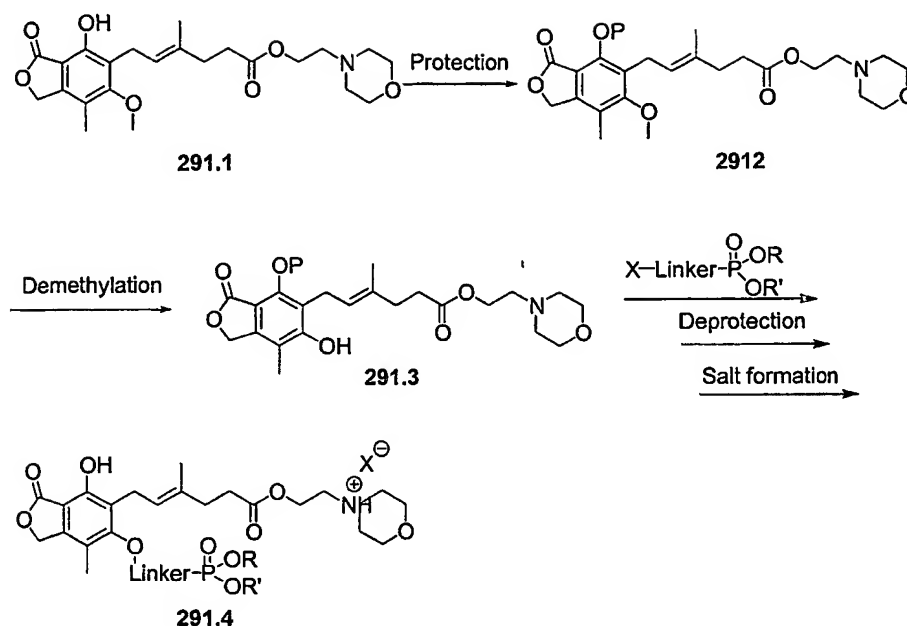
Example 290

Representative compounds of the invention can be prepared as illustrated above. The carboxylic acid of mycophenolic acid can be replaced with a phosphonic acid that may also serve as a prodrug handle. In order to remove the carboxylic acid containing side chain, the acid chloride 290.5 (prepared in Example 289) is converted to ester 290.1. Protection of the phenol with a silyl group, followed by dihydroxylation and cleavage of the diol generates aldehyde 290.3 (Pankiewicz, et al., *J. Med. Chem.*, 2002, 45, 703), (Patterson et al., U.S. Patent No. 5,444,072). A Wittig reaction with ylide 290.4 carrying an appropriately protected phosphonate provides the desired compound 290.5. Final deprotection yields compound 290.6. A specific compound of the invention can be prepared as follows.

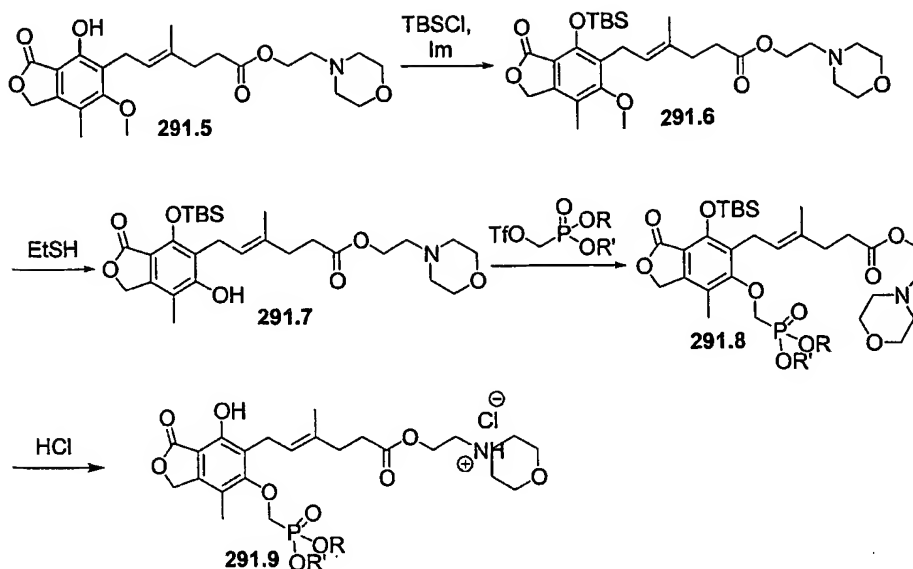


Mycophenolate ester **290.8** can simply be prepared by stirring the acid chloride **290.7** with MeOH. Then, the phenol position of mycophenolate ester is protected by a silyl group such as TBS to provide compound **290.9**. Once the phenol position is protected, dihydroxylation using osmium tetroxide followed by periodate cleavage provides aldehyde **290.10**. Aldehyde **290.10** and excess of the ylide **290.11** are heated in benzene at reflux for about 24 hours. The reaction mixture is concentrated and the residue is purified by column chromatography to provide olefin **290.12** (Pankiewics et al., *J. Med. Chem.*, 2002, 45, 703). A final deprotection using HF-pyridine yields the final product **290.13**.

Example 291

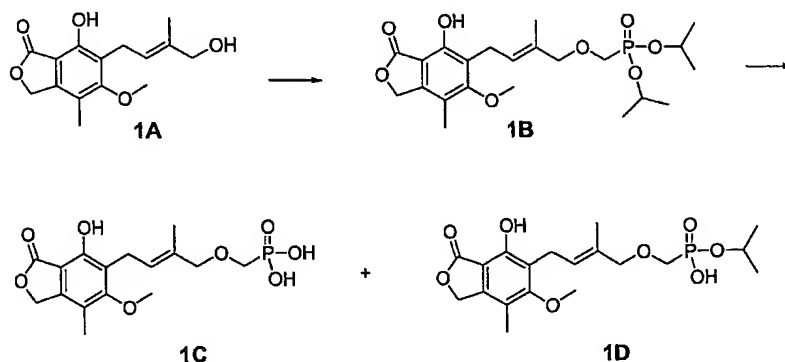


Representative compounds of the invention can be prepared as illustrated above. Another attachment point of the compound can be unmasked after demethylation of mycophenolate ester **291.2** as illustrated above. For this purpose, the 4-OH needs to be masked with a protecting group (P) such as a silyl group. Once the 6-MeO is demethylated and alkylated, the protecting group at position 4 is removed to reveal the final product **291.4**. The morphonyl ethanol group is installed early and carried through the alkylation steps. A different protecting group may be installed initially and removed later. In such the latter type of synthesis, the last step is the formation of the morpholinoethyl ester prodrug. A specific compound of the invention can be prepared as described below.



Phenol **291.5** is protected with TBS group in CH_2Cl_2 using imidazole as base to yield **291.6**. Demethylation is performed using thiolate nucleophiles to generate compound **291.7**. A variety of other methods are also available in literature as described in *Protective Groups in Organic Synthesis* by Greene and Wuts. Alkylation of the 6-OH using a triflate of the phosphonate proceeds well using K_2CO_3 or TEA to provide **291.8**. Final deprotection to remove the TBS group provides product **291.9**.

Example 292



Representative compounds of the invention can be prepared as illustrated above.

[4-(4-Hydroxy-6-methoxy-7-methyl-3-oxo-1,3-dihydro-isobenzofuran-5-yl)-2-methyl-but-2-enyloxymethyl]-phosphonic acid diisopropyl ester

A mixture of 7-hydroxy-6-(4-hydroxy-3-methyl-but-2-enyl)-5-methoxy-4-methyl-3H-isobenzofuran-1-one 1A (50 mg, 0.18 mmol, Pankiewicz *et al.*, *J. Med. Chem.*, **45**, 703), diisopropyl bromomethylphosphonate (93 mg, 0.36 mmol) and lithium *t*-butoxide (1M in THF, 0.54 mL) in DMF (3 mL) was heated at 70 °C for 5 hours. The reaction was quenched with 1N HCl. The mixture was poured into 5 % aqueous lithium chloride, extracted with ethyl acetate, and concentrated. The residue was purified by chromatography on silica gel, affording [4-(4-hydroxy-6-methoxy-7-methyl-3-oxo-1,3-dihydro-isobenzofuran-5-yl)-2-methyl-but-2-enyloxymethyl]-phosphonic acid diisopropyl ester 1B (25 mg, 32%); ¹H NMR (300 MHz, CDCl₃) δ 1.25 (m, 12H), 1.79 (s, 3H), 2.05 (s, 3H), 3.37 (d, *J* = 6.6 Hz, 2H), 3.58 (d, 2H), 3.77 (s, 3H), 3.97 (m, 2H), 4.68 (m, 2H), 5.19 (s, 2H), 5.45 (t, *J* = 6.6 Hz, 1H), 7.83 (s, 1H) ppm.

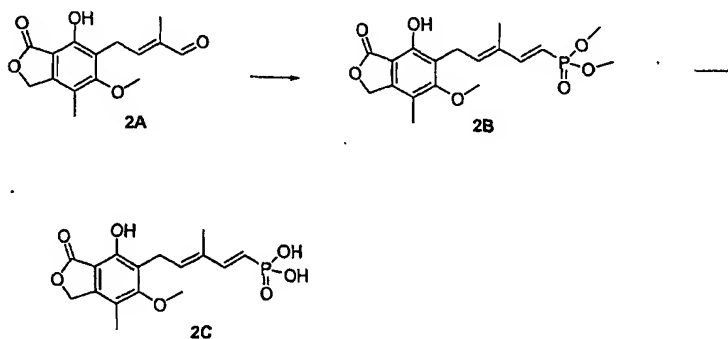
[4-(4-Hydroxy-6-methoxy-7-methyl-3-oxo-1,3-dihydro-isobenzofuran-5-yl)-2-methyl-but-2-enyloxymethyl]-phosphonic acid and [4-(4-Hydroxy-6-methoxy-7-methyl-3-oxo-1,3-dihydro-isobenzofuran-5-yl)-2-methyl-but-2-enyloxymethyl]-phosphonic acid monoisopropyl ester

To a solution of [4-(4-hydroxy-6-methoxy-7-methyl-3-oxo-1,3-dihydro-isobenzofuran-5-yl)-2-methyl-but-2-enyloxymethyl]-phosphonic acid diisopropyl ester 1B (25 mg, 0.055 mmol) and 2,6-lutidine (0.18 mL, 1.65 mmol) in acetonitrile was added trimethylsilyl bromide (0.126 mL, 1.1 mmol) at 0°C. The mixture was allowed to warm to room temperature and stirred for 4 hours. The reaction was quenched with methanol at 0°C, and the resulting mixture was concentrated. The residue was purified by preparative reverse-phase HPLC to afford, after removal of the solvent, [4-(4-hydroxy-6-methoxy-7-methyl-3-oxo-1,3-dihydro-isobenzofuran-5-yl)-2-methyl-but-2-enyloxymethyl]-phosphonic acid 1C as an oil (17 mg, 83 %); ¹H NMR (300 MHz, CD₃OD) δ 1.81 (s, 3H), 2.06 (s, 3H), 3.40 (d, *J* = 6.6 Hz, 2H), 3.50 (d, 2H), 3.77 (s, 3H), 3.97 (s, 2H), 5.20 (s, 2H), 5.47 (t, *J* = 6.6 Hz, 1H) and [4-(4-hydroxy-6-methoxy-7-methyl-3-oxo-1,3-dihydro-isobenzofuran-5-yl)-2-methyl-but-2-enyloxymethyl]-phosphonic acid monoisopropyl ester 1D as an oil (2 mg, 7 %);

^1H NMR (300 MHz, CD_3OD) δ 1.23 (d, 6H), 1.81 (s, 3H), 2.08 (s, 3H), 3.40 (d, $J = 6.6$ Hz, 2H), 3.50 (d, 2H), 3.77 (s, 3H), 3.90 (s, 2H), 4.50 (m, 1H), 5.20 (s, 2H), 5.47 (t, $J = 6.6$ Hz, 1H) ppm.

Example 293

Representative compounds of the invention can be prepared as illustrated below.



[5-(4-Hydroxy-6-methoxy-7-methyl-3-oxo-1,3-dihydro-isobenzofuran-5-yl)-3-methyl-penta-1,3-dienyl]-phosphonic acid dimethyl ester

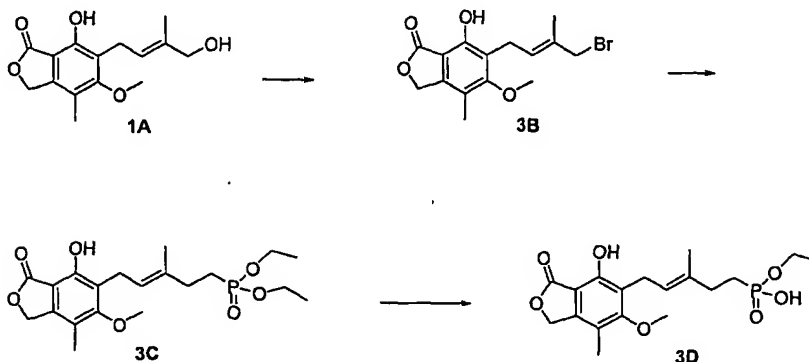
To a solution of tetramethylmethylene diphosphonate (102 mg, 0.44 mmol) in THF (2.5 mL) was added a THF solution of sodium bis(trimethylsilyl)amide (1.0 M, 0.44 mL). After stirring for 30 minutes, a solution of 4-(4-hydroxy-6-methoxy-7-methyl-3-oxo-1,3-dihydro-isobenzofuran-5-yl)-2-methylbut-2-enal 2A (30 mg, 0.11 mmol, Pankiewicz *et al.*, *J. Med. Chem.*, 45, 703) in THF (2.5 mL) was added, and stirring was continued for an additional 15 minutes. The reaction was quenched with saturated aqueous ammonium chloride. The mixture was extracted with ethyl acetate. After evaporation of solvent, the residue was purified by chromatography on silica gel eluting with ethyl acetate (50 % to 100 %) / hexanes, affording [5-(4-hydroxy-6-methoxy-7-methyl-3-oxo-1,3-dihydro-isobenzofuran-5-yl)-3-methyl-penta-1,3-dienyl]-phosphonic acid dimethyl ester 2B (30 mg, 71 %) as an oil; ^1H NMR (300 MHz, CDCl_3) δ 1.80 (s, 3H), 2.04 (s, 3H), 3.45 (d, $J = 6.6$ Hz, 2H), 3.76 (s, 3H), 3.88 (d, 6H), 5.20 (s, 3H), 5.55 (m, 1H), 5.95 (m, 1H), 7.05 (m, 1H), 7.65 (s, 1H) ppm.

[5-(4-Hydroxy-6-methoxy-7-methyl-3-oxo-1,3-dihydro-isobenzofuran-5-yl)-3-methyl-penta-1,3-dienyl]-phosphonic acid

To a solution of [5-(4-hydroxy-6-methoxy-7-methyl-3-oxo-1,3-dihydro-isobenzofuran-5-yl)-3-methyl-penta-1,3-dienyl]-phosphonic acid dimethyl ester 2B (22 mg, 0.057 mmol) and 2,6-lutidine (0.22 mL, 1.71 mmol) in acetonitrile was added trimethylsilyl bromide (0.183 mL, 1.71 mmol) at 0°C. The mixture was allowed to warm to room temperature and stirred for 1 hour. The reaction was quenched with methanol at 0°C, and the resulting mixture was concentrated. The residue was purified by preparative reverse-phase HPLC to afford, after removal of the solvent, [5-(4-hydroxy-6-methoxy-7-methyl-3-oxo-1,3-dihydro-isobenzofuran-5-yl)-3-methyl-penta-1,3-dienyl]-phosphonic acid 2C as a solid (13 mg, 65 %); ¹H NMR (300 MHz, CD₃OD) δ 1.91 (s, 3H), 2.10 (s, 3H), 3.55 (d, *J* = 6.6 Hz, 2H), 3.75 (s, 3H), 5.2 (s, 2H), 5.6-5.8 (m, 2H), 6.9 (m, 1H) ppm.

Example 294

Representative compounds of the invention can be prepared as illustrated below.



6-(4-Bromo-3-methyl-but-2-enyl)-7-hydroxy-5-methoxy-4-methyl-3H-isobenzofuran-1-one

Polymer-supported triphenylphosphine (3 mmol/g, 0.5 g) was soaked in dichloromethane (10mL) for 1 hour. 7-Hydroxy-6-(4-hydroxy-3-methyl-but-2-enyl)-5-methoxy-4-methyl-3H-isobenzofuran-1-one 1A (100 mg, 0.36 mmol) and carbon tetrabromide (143 mg, 0.43 mmol) were sequentially added and the

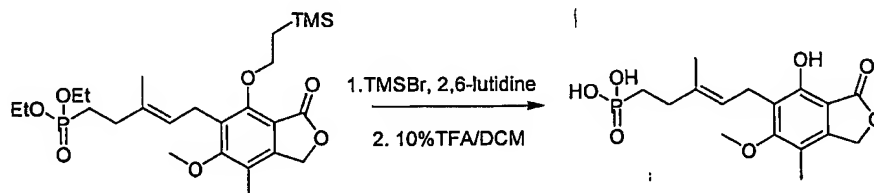
mixture was shaken for 1 hour at room temperature. More carbon tetrabromide (143 mg, 0.43 mmol) was added and the mixture was shaken further for 1 hour. The mixture was filtered and the filtrate was concentrated. The residue was chromatographed on silica gel (0 % to 60 % ethyl acetate / hexanes) to afford 6-(4-bromo-3-methyl-but-2-enyl)-7-hydroxy-5-methoxy-4-methyl-3H-isobenzofuran-1-one 3B as an oil (52 mg, 42 %); ^1H NMR (300 MHz, CDCl_3) δ 1.95 (s, 3H), 2.16 (s, 3H), 3.44 (d, $J = 7.2$ Hz, 2H), 3.78 (s, 3H), 3.98 (s, 2H), 5.21 (s, 2H), 5.68 (t, $J = 7.2$ Hz, 1H), 7.71 (brs, 1H) ppm.

[5-(4-Hydroxy-6-methoxy-7-methyl-3-oxo-1,3-dihydro-isobenzofuran-5-yl)-3-methyl-pent-3-enyl]-phosphonic acid diethyl ester

n-Butyl lithium (1.6 M in hexanes, 1 mL) was added to an equal volume of THF at -20°C . A solution of diethyl methylphosphonate (220 mg, 1.45 mmol) in THF (1 mL) was then added dropwise and the solution was stirred for 30 minutes. After cooling at -60°C , the solution was transferred *via* a cannula to a vial containing copper (I) iodide (276 mg, 1.45 mmol), and the resulting mixture was stirred for 1 hour at -30°C . A solution of 6-(4-bromo-3-methyl-but-2-enyl)-7-hydroxy-5-methoxy-4-methyl-3H-isobenzofuran-1-one 3B (50 mg, 0.15 mmol) in THF (1 mL) was added and the mixture was allowed to warm to 0°C for 2 hours before saturated aqueous ammonium chloride was added. The reaction mixture was acidified with 2 N HCl and extracted with ethyl acetate. The ethyl acetate extract was concentrated and the residue was chromatographed on silica gel (40% to 100% ethyl acetate / hexanes), affording [5-(4-hydroxy-6-methoxy-7-methyl-3-oxo-1,3-dihydro-isobenzofuran-5-yl)-3-methyl-pent-3-enyl]-phosphonic acid diethyl ester 3C as an oil (27 mg, contaminated with the starting diethyl methylphosphonate); ^1H NMR (300 MHz, CDCl_3) δ 1.32 (m, 6H), 1.8-1.9 (m, 5H), 2.18 (s, 3H), 2.25 (m, 2H), 3.42 (d, $J = 7.2$ Hz, 2H), 3.78 (s, 3H), 4.15 (m, 4H), 5.21 (s, 2H), 5.24 (t, $J = 7.2$ Hz, 1H), 7.65 (s, 1H) ppm.

[5-(4-Hydroxy-6-methoxy-7-methyl-3-oxo-1,3-dihydro-isobenzofuran-5-yl)-3-methyl-pent-3-enyl]-phosphonic acid monoethyl ester

A mixture of [5-(4-hydroxy-6-methoxy-7-methyl-3-oxo-1,3-dihydro-isobenzofuran-5-yl)-3-methyl-pent-3-enyl]-phosphonic acid diethyl ester 3C (27 mg, 0.066 mmol), LiOH (200 mg), MeOH (3 mL) and water (1 mL) was stirred at 70°C for 4 hours. After cooling, the reaction solution was acidified with 2 N HCl, mixed with brine, and extracted with ethyl acetate / acetonitrile. The organic extract was concentrated and the residue was purified by preparative reverse-phase HPLC (acetonitrile and 0.1% aqueous CF₃COOH), affording [5-(4-hydroxy-6-methoxy-7-methyl-3-oxo-1,3-dihydro-isobenzofuran-5-yl)-3-methyl-pent-3-enyl]-phosphonic acid monoethyl ester 3D (7 mg, 28 %); ¹H NMR (300 MHz, CD₃OD) δ 1.28 (t, *J* = 6.9 Hz, 3H), 1.7-1.9 (m, 5H), 2.20 (s, 3H), 2.2-2.3 (m, 2H), 3.41 (d, *J* = 6.6 Hz, 2H), 3.80 (s, 3H), 4.02 (m, 2H), 5.2-5.3 (m, 3H) ppm.



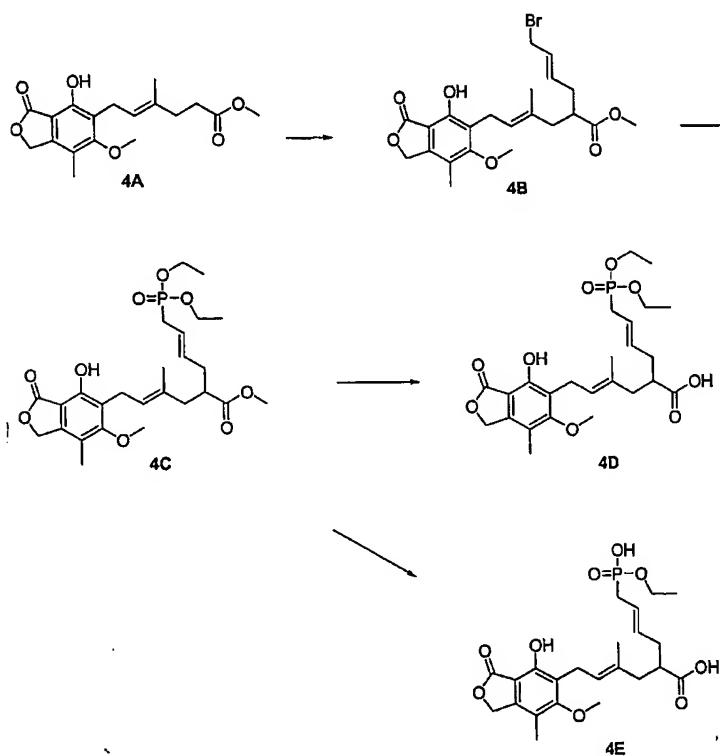
[5-(4-Hydroxy-6-methoxy-7-methyl-3-oxo-1,3-dihydro-isobenzofuran-5-yl)-3-methyl-pent-3-enyl]-phosphonic acid

To a solution of {5-[6-Methoxy-7-methyl-3-oxo-4-(2-trimethylsilyl-ethoxy)-1,3-dihydro-isobenzofuran-5-yl]-3-methyl-pent-3-enyl}-phosphonic acid diethyl ester (20 mg, 0.039 mmol) in DMF (0.5 mL) and DCM (0.5 mL) was added TMSBr (50.5 μL, 0.39 mmol) followed by 2,6-lutidine (45.3 μL, 0.39 mmol). The reaction was allowed to proceed for one hour when it was complete, as judged by LCMS. The reaction mixture was quenched with MeOH and concentrated to dryness. The residue was purified by preparative reverse-phase HPLC. The fraction containing the desired product was concentrated and treated with 10% TFA/DCM for 5 minutes. After concentration, the residue was purified by preparative reverse-phase HPLC to provide 7 mg (50%) of [5-(4-Hydroxy-6-methoxy-7-methyl-3-oxo-1,3-dihydro-isobenzofuran-5-yl)-3-methyl-pent-3-

enyl]-phosphonic acid as a solid. ^1H NMR (300 MHz, CD_3OD) δ 1.66-1.78 (m, 5H), 2.10 (s, 3H), 2.16-2.22 (m, 2H), 3.34 (d, $J = 7.2$ Hz, 2H), 3.72 (s, 3H), 5.16 (s, 2H), 5.20 (t, $J = 7.2$ Hz, 1H) ppm; ^{31}P (121.4 MHz, CD_3OD) δ 31.57 ppm; MS (m/z) 355 $[\text{M}-\text{H}]^-$, 357 $[\text{M}+\text{H}]^+$.

Example 295

Representative compounds of the invention can be prepared as illustrated below.



2-(4-Bromo-but-2-enyl)-6-(4-hydroxy-6-methoxy-7-methyl-3-oxo-1,3-dihydro-isobenzofuran-5-yl)-4-methyl-hex-4-enoic acid methyl ester

To a cooled (-78°C) solution of mycophenolic acid methyl ester 4A (138 mg, 0.41 mmol) in THF (2.5 mL) was added a THF solution of sodium bis(trimethylsilyl)amide (1.0 M, 0.98 mL). After stirring for 30 minutes, a solution of 1,4-dibromo-2-butene (950 mg, 4.1 mmol) in THF (2.5 mL) was added and stirring was continued for 10 minutes. The resulting mixture was warmed to -30°C and stored at this temperature for 16 hours. The reaction was

quenched with saturated aqueous ammonium chloride. The mixture was extracted with ethyl acetate to give, after evaporation of the solvent, a residue that was purified by chromatography on silica gel eluting with ethyl acetate (0 % to 40 %) / hexanes, affording 2-(4-bromo-but-2-enyl)-6-(4-hydroxy-6-methoxy-7-methyl-3-oxo-1,3-dihydro-isobenzofuran-5-yl)-4-methyl-hex-4-enoic acid methyl ester **4B** (150 mg, 78 %) as an oil; ^1H NMR (300 MHz, CDCl_3) δ 1.75 (s, 3H), 2.0-2.4 (m, 8H), 2.62 (m, 1H), 3.37 (d, $J = 6.6$ Hz, 2H), 3.58 (s, 3H), 3.76 (s, 3H), 3.88 (d, $J = 4.8$ Hz, 2H), 5.1-5.3 (m, 3H), 5.67 (brs, 2H), 7.67 (s, 1H) ppm.

2-[4-(Diethoxy-phosphoryl)-but-2-enyl]-6-(4-hydroxy-6-methoxy-7-methyl-3-oxo-1,3-dihydro-isobenzofuran-5-yl)-4-methyl-hex-4-enoic acid methyl ester

A solution of 2-(4-bromo-but-2-enyl)-6-(4-hydroxy-6-methoxy-7-methyl-3-oxo-1,3-dihydro-isobenzofuran-5-yl)-4-methyl-hex-4-enoic acid methyl ester **4B** (140 mg, 0.30 mmol) and triethylphosphite (600 mg, 3.6 mmol) in toluene (30 mL) was stirred at reflux for 20 hours. The mixture was concentrated and chromatographed on silica gel eluting with ethyl acetate (60 % to 100 %) / hexanes, affording 2-[4-(diethoxy-phosphoryl)-but-2-enyl]-6-(4-hydroxy-6-methoxy-7-methyl-3-oxo-1,3-dihydro-isobenzofuran-5-yl)-4-methyl-hex-4-enoic acid methyl ester **4C** as an oil (70 mg, 43 %); ^1H NMR (300 MHz, CDCl_3) δ 1.27 (m, 6H), 1.79 (s, 3H), 2.0-2.7 (m, 8H), 3.37 (d, $J = 6.6$ Hz), 3.52 (s, 3H), 3.75 (s, 3H), 4.08 (m, 4H), 5.20 m, 3H), 5.45 (m, 2H) ppm.

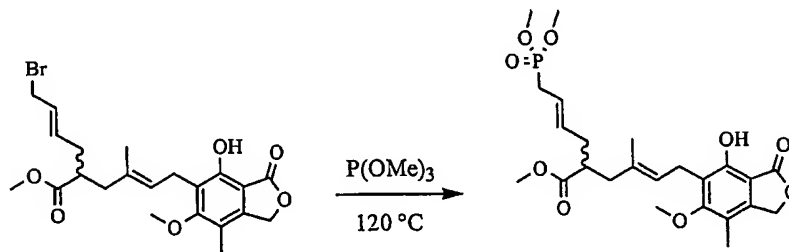
2-[4-(Diethoxy-phosphoryl)-but-2-enyl]-6-(4-hydroxy-6-methoxy-7-methyl-3-oxo-1,3-dihydro-isobenzofuran-5-yl)-4-methyl-hex-4-enoic acid

A mixture of 2-[4-(diethoxy-phosphoryl)-but-2-enyl]-6-(4-hydroxy-6-methoxy-7-methyl-3-oxo-1,3-dihydro-isobenzofuran-5-yl)-4-methyl-hex-4-enoic acid methyl ester **4C** (33 mg, 0.063 mmol) and lithium hydroxide (44 mg) in a mixture of THF (6 mL) and water (1 mL) was stirred at room temperature for 6 hours. The organic solvent was removed and the residue was partitioned between ethyl acetate and 5 % aqueous sodium bicarbonate. The aqueous layer

was acidified with 2 N HCl and extracted with ethyl acetate. The ethyl acetate extract was concentrated, affording 2-[4-(diethoxy-phosphoryl)-but-2-enyl]-6-(4-hydroxy-6-methoxy-7-methyl-3-oxo-1,3-dihydro-isobenzofuran-5-yl)-4-methyl-hex-4-enoic acid **4D** as an oil (30 mg, 100%); ^1H NMR (300 MHz, CDCl_3) δ 1.27 (m, 6H), 1.79 (s, 3H), 2.0-2.7 (m, 8H), 3.37 (d, $J = 6.6$ Hz), 3.75 (s, 3H), 4.08 (m, 4H), 5.19 (s, 2H), 5.25 (m, 1H), 5.44 (m, 1H), 5.55 (m, 1H), 5.45 (m, 2H) ppm.

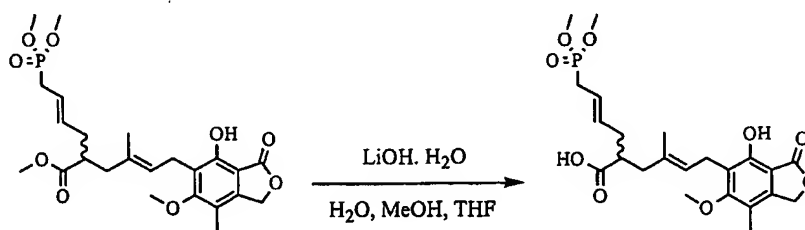
2-[4-(Ethoxy-hydroxy-phosphoryl)-but-2-enyl]-6-(4-hydroxy-6-methoxy-7-methyl-3-oxo-1,3-dihydro-isobenzofuran-5-yl)-4-methyl-hex-4-enoic acid

A mixture of 2-[4-(diethoxy-phosphoryl)-but-2-enyl]-6-(4-hydroxy-6-methoxy-7-methyl-3-oxo-1,3-dihydro-isobenzofuran-5-yl)-4-methyl-hex-4-enoic acid methyl ester **4C** (25 mg, 0.048 mmol) and lithium hydroxide (200 mg) in a mixture of methanol (3 mL) and water (1 mL) was stirred at 70°C for 2 hours. The organic solvent was evaporated and the residue acidified with 2N HCl and extracted with ethyl acetate /acetonitrile. The organic extract was concentrated, and the residue was purified by preparative reverse-phase HPLC (acetonitrile and 0.1% aqueous CF_3COOH), affording 2-[4-(ethoxy-hydroxy-phosphoryl)-but-2-enyl]-6-(4-hydroxy-6-methoxy-7-methyl-3-oxo-1,3-dihydro-isobenzofuran-5-yl)-4-methyl-hex-4-enoic acid **4E** as an oil (15 mg, 89%); ^1H NMR (300 MHz, CD_3OD) δ 1.25 (t, $J = 6.9$ Hz, 3H), 1.81 (s, 3H), 2.1-2.6 (m, 8H), 3.40 (d, $J = 6.6$ Hz, 2H), 3.77 (s, 3H), 3.97 (m, 2H), 5.1-5.3 (m, 3H), 5.67 (brs, 2H) ppm.



2-[4-(Dimethoxy-phosphoryl)-but-2-enyl]-6-(4-hydroxy-6-methoxy-7-methyl-3-oxo-1,3-dihydro-isobenzofuran-5-yl)-4-methyl-hex-4-enoic acid methyl ester

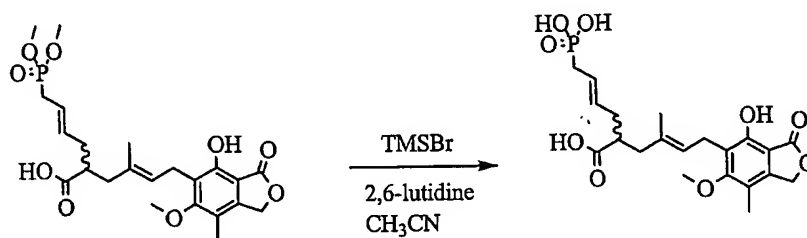
Under a N_2 atmosphere, a solution of 2-(4-bromo-but-2-enyl)-6-(4-hydroxy-6-methoxy-7-methyl-3-oxo-1,3-dihydro-isobenzofuran-5-yl)-4-methyl-hex-4-enoic acid methyl ester (490 mg, 1.05 mmol) in trimethylphosphite (2.5 mL, 21.1 mmol) was heated at 120°C for 1 hour. The reaction was allowed to cool to room temperature. The reaction mixture was worked up by removal of the solvent *in vacuo* followed by chromatography using EtOAc-hexanes to provide 460 mg (88%) of the product as an oil. 1H NMR (300 MHz, $CDCl_3$) δ 1.77 (s, 3H), 2.081- 2.31 (m, 4H), 2.15 (s, 3H), 2.52 (d, 1H, $J = 22$ Hz), 2.54 (d, 1H, $J = 22$ Hz), 2.55- 2.63 (m, 1H), 3.36 (d, 2H, $J = 7$ Hz), 3.57 (s, 3H), 3.72 (d, 6H, $J = 11$ Hz), 3.76 (s, 3H), 5.20 (s, 2H), 5.20- 5.26 (m, 1H), 5.36- 5.56 (m, 2H), 7.69 (s, 1H) ppm; ^{31}P (121.4 MHz, $CDCl_3$) δ 30.1 ppm; MS (m/z) 497.2 $[M+H]^+$, 519.2 $[M+Na]^+$.



2-[4-(Dimethoxy-phosphoryl)-but-2-enyl]-6-(4-hydroxy-6-methoxy-7-methyl-3-oxo-1,3-dihydro-isobenzofuran-5-yl)-4-methyl-hex-4-enoic acid

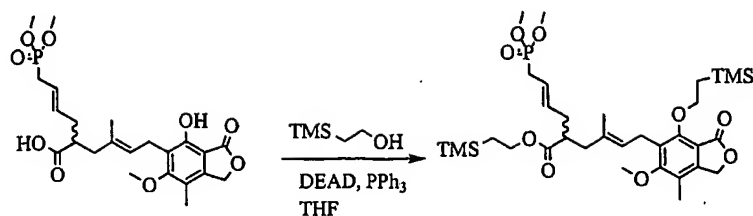
2-[4-(Dimethoxy-phosphoryl)-but-2-enyl]-6-(4-hydroxy-6-methoxy-7-methyl-3-oxo-1,3-dihydro-isobenzofuran-5-yl)-4-methyl-hex-4-enoic acid methyl ester (460 mg, 0.927 mmol) in a solution of 1: 1: 2 of H_2O , MeOH, THF (8 mL) was stirred with $LiOH \cdot H_2O$ (78 mg, 1.86 mmol) at ambient temperature for 12 hours. A second batch of $LiOH \cdot H_2O$ (40 mg, 0.952 mmol) was added. The reaction mixture was stirred at room temperature for another 16 hours, after which no further progress was observed. The reaction was quenched by addition of a saturated aqueous solution of NH_4Cl . The organic layer was removed *in*

vacuo and the product was extracted with EtOAc from the aqueous layer, which had been acidified by addition of 5 drops of 2 N HCl. The product was further purified by chromatography to provide the desired product. ^1H NMR (300 MHz, CDCl_3) δ 1.79 (s, 3H), 2.08- 2.38 (m, 4H), 2.15 (s, 3H), 2.53 (d, 1H, $J=22$ Hz), 2.60 (d, 1H, $J=22$ Hz), 2.57- 2.64 (m, 1H), 3.38 (d, 2H, $J=7$ Hz), 3.72 (d, 6H, $J=11$ Hz) 3.76 (s, 3H), 5.20 (s, 2H), 5.27 (t, 1H, $J=6$ Hz), 5.36- 5.63 (m, 2H) ppm; ^{31}P (121.4 MHz, CDCl_3) δ 30.5 ppm; MS (m/z) 481.2 $[\text{M}-\text{H}]^-$.



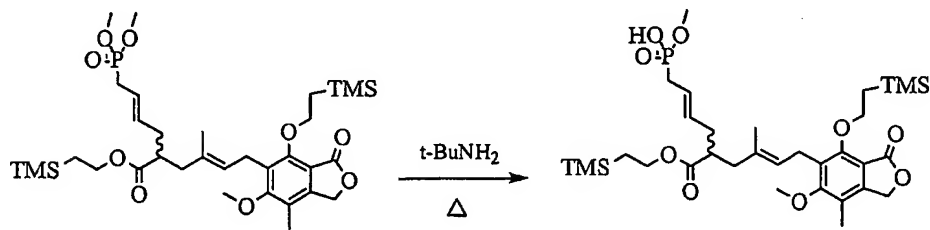
2-[4-(2-[4-(Dimethoxy-phosphoryl)-but-2-enyl]-6-(4-hydroxy-6-methoxy-7-methyl-3-oxo-1,3-dihydro-isobenzofuran-5-yl)-4-methyl-hex-4-enoic acid

To a solution of 2-[4-(dimethoxy-phosphoryl)-but-2-enyl]-6-(4-hydroxy-6-methoxy-7-methyl-3-oxo-1,3-dihydro-isobenzofuran-5-yl)-4-methyl-hex-4-enoic acid (25 mg, 0.052 mmol) in acetonitrile (2 mL) was added 2,6-lutidine (60 μL , 0.52 mmol) and TMSBr (67 μL , 0.52 mmol). The reaction was allowed to proceed for 45 minutes when it was completed as judged by LCMS. The reaction mixture was concentrated under reduced pressure and quenched with an aqueous NaOH solution (1 mL). The product was purified by RP HPLC (using a C18 column with a gradient of H_2O , 0.1% TFA-acetonitrile, 0.1% TFA) to provide 14.2 mg (60%) of the product as a solid. ^1H NMR (300 MHz, CD_3OD) δ 1.81 (s, 3H), 2.081- 2.31 (m, 4H), 2.16 (s, 3H), 2.45 (d, 1H, $J=22$ Hz), 2.47 (d, 1H, $J=22$ Hz), 2.55- 2.63 (m, 1H), 3.38 (d, 2H, $J=7$ Hz), 3.77 (s, 3H), 5.25 (s, 2H), 5.20- 5.36 (m, 1H), 5.36- 5.56 (m, 2H) ppm; ^{31}P (121.4 MHz, CD_3OD) δ 25.4 ppm; MS (m/z) 453 $[\text{M}-\text{H}]^-$.



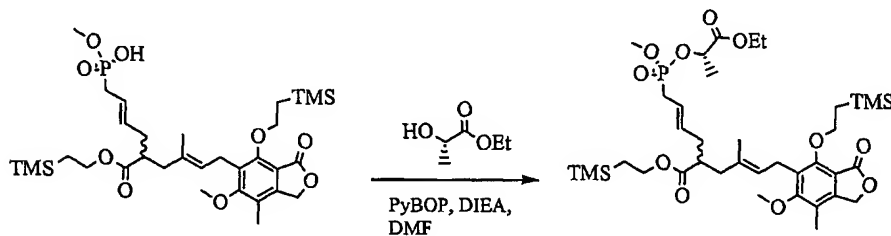
2-[4-(Dimethoxy-phosphoryl)-but-2-enyl]-6-[6-methoxy-7-methyl-3-oxo-4-(2-trimethylsilyl-ethoxy)-1,3-dihydro-isobenzofuran-5-yl]-4-methyl-hex-4-enoic acid 2-trimethylsilyl-ethyl ester

A solution of 2-[4-(dimethoxy-phosphoryl)-but-2-enyl]-6-(4-hydroxy-6-methoxy-7-methyl-3-oxo-1,3-dihydro-isobenzofuran-5-yl)-4-methyl-hex-4-enoic acid (160 mg, 0.332 mmol) and trimethylsilylethanol (160 mg, 1.36 mmol) in THF (8.00 mL) was stirred with triphenylphosphine (345 mg, 1.33 mmol). To this solution was added diethyl azodicarboxylate (230 μ L, 1.33 mmol) at 0°C. The mixture was allowed to warm to room temperature and stirred for 16 hours. Additional triphenylphosphine (180 mg, 0.692 mmol), trimethylsilylethanol (160 mg, 1.36 mmol), and diethyl azodicarboxylate (115 μ L, 0.665 mmol) were added and the reaction mixture was stirred for another 1 day at room temperature. The reaction was worked up by removing the solvents *in vacuo* and purifying the residue by silica gel chromatography to provide 192 mg (85%) of the product as a clear oil. ^1H NMR (300 MHz, CDCl_3) δ 0.03 (s, 9H), 0.05 (s, 9H), 0.93- 0.96 (m, 2H), 1.20- 1.29 (m, 2H), 1.78 (s, 3H), 2.01- 2.32 (m, 4H), 2.17 (s, 3H), 2.51 (d, 1H, $J=22$ Hz), 2.58 (d, 1H, $J=22$ Hz), 2.50- 2.60 (m, 1H), 3.37 (d, 2H, $J=7$ Hz), 3.72 (d, 6H, $J=11$ Hz), 3.76 (s, 3H), 4.08 (appt t, 2H, $J=8$ Hz), 4.30 (appt t, 2H, $J=8$ Hz), 5.12 (s, 2H), 5.15- 5.25 (m, 1H), 5.36- 5.63 (m, 2H) ppm; ^{31}P (121.4 MHz, CDCl_3) δ 29.3 ppm; MS (m/z) 705.3 $[\text{M}+\text{Na}]^+$.



2-[4-(Hydroxy-methoxy-phosphoryl)-but-2-enyl]-6-[6-methoxy-7-methyl-3-oxo-4-(2-trimethylsilylanyl-ethoxy)-1,3-dihydro-isobenzofuran-5-yl]-4-methyl-hex-4-enoic acid 2-trimethylsilylanyl-ethyl ester

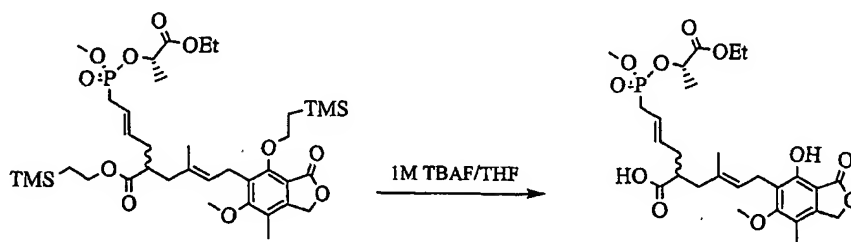
A mixture of 2-[4-(dimethoxy-phosphoryl)-but-2-enyl]-6-[6-methoxy-7-methyl-3-oxo-4-(2-trimethylsilylanyl-ethoxy)-1,3-dihydro-isobenzofuran-5-yl]-4-methyl-hex-4-enoic acid 2-trimethylsilylanyl-ethyl ester (184 mg, 0.270 mmol) in *tert*-butylamine (2.8 mL, 27 mmol) was heated at 60 °C for 24 hours. The solution was allowed to cool to room temperature and concentrated. The residue was purified by silica gel column chromatography using MeOH/ CH₂Cl₂ (0-30%) to provide 75 mg of the product as a clear oil. ¹H NMR (300 MHz, CDCl₃) δ 0.01 (s, 9H), 0.04 (s, 9H), 0.89 (appt t, 2H, *J*= 9 Hz), 1.23 (appt t, 2H, *J*= 9 Hz), 1.77 (s, 3H), 2.01- 2.31 (m, 4H), 2.17 (s, 3H), 2.36 (d, 1H, *J*= 22 Hz), 2.38 (d, 1H, *J*= 22 Hz), 2.52 (septet, 1H, *J*= 9 Hz), 3.39 (d, 2H, *J*= 7 Hz), 3.51 (d, 3H, *J*= 11 Hz), 4.01- 4.08 (m, 2H), 4.30 (dd, 2H, *J*= 8, 9 Hz), 5.11 (s, 2H), 5.19 (br t, 1H, *J*= 6 Hz), 5.33- 5.56 (m, 2H), 8.49 (br s, 1H) ppm; ³¹P (121.4 MHz, CDCl₃) δ 22.1 ppm; MS (*m/z*) 667.4 [M+Na]⁺.



2-[4-[(1-Ethoxycarbonyl-ethoxy)-methoxy-phosphoryl]-but-2-enyl]-6-[6-methoxy-7-methyl-3-oxo-4-(2-trimethylsilylanyl-ethoxy)-1,3-dihydro-isobenzofuran-5-yl]-4-methyl-hex-4-enoic acid 2-trimethylsilylanyl-ethyl ester

A solution of 2-[4-(hydroxy-methoxy-phosphoryl)-but-2-enyl]-6-[6-methoxy-7-methyl-3-oxo-4-(2-trimethylsilylanyl-ethoxy)-1,3-dihydro-isobenzofuran-5-yl]-4-methyl-hex-4-enoic acid 2-trimethylsilylanyl-ethyl ester (67 mg, 0.10 mmol) and PyBOP (234 mg, 0.450 mmol) in DMF (1.5 mL) was stirred with ethyl (S)-(-)-lactate (53 mg, 0.45 mmol) and DIEA (174 μL, 1.00 mmol) at ambient temperature for 1 hour, when complete consumption of the

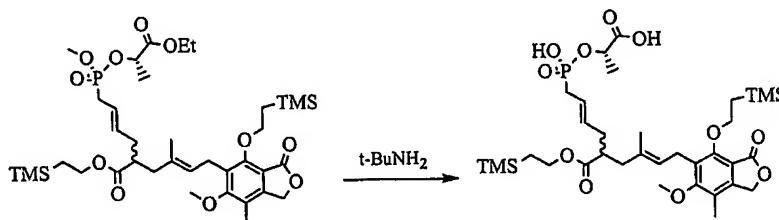
starting materials was observed. The reaction was worked up by addition of saturated aqueous sodium chloride and ethyl acetate. The organic layer was separated and washed with 5% aqueous solution of lithium chloride. The organic layer was dried *in vacuo* and the residue was purified by silica gel chromatography using MeOH-CH₂Cl₂ (0-20%) to provide 57 mg (74%) of the desired product as a clear oil. ¹H NMR (300 MHz, CDCl₃) δ 0.02 (s, 9H), 0.05 (s, 9H), 0.88- 0.94 (m, 2H), 1.20- 1.30 (m, 2H), 1.29 (t, 3H, *J*= 7 Hz), 1.45 (d, 3H, *J*= 7 Hz), 1.78 (s, 3H), 2.01- 2.31 (m, 4H), 2.17 (s, 3H), 2.50- 2.58 (m, 1H), 2.65 (d, 1H, *J*= 22 Hz), 2.67 (d, 1H, *J*= 22 Hz), 3.39 (d, 2H, *J*= 7 Hz), 3.69 and 3.77 (d, 3H, *J*= 11 Hz), 3.76 (s, 3H), 4.07 (appt t, 2H, *J*= 7 Hz), 4.20 (dq, 2H, *J*= 3, 7 Hz), 4.29 (appt t, 2H, *J*= 9 Hz), 4.85- 4.99 (m, 1H), 5.12 (s, 2H), 5.19 (br t, 1H, *J*= 6 Hz), 5.33- 5.61 (m, 2H) ppm; ³¹P (121.4 MHz, CDCl₃) δ 28.9, 29.9 ppm; MS (*m/z*) 791.4 [M+Na]⁺.



2-{4-[(1-Ethoxycarbonyl-ethoxy)-methoxy-phosphoryl]-but-2-enyl}-6-(4-hydroxy-6-methoxy-7-methyl-3-oxo-1,3-dihydro-isobenzofuran-5-yl)-4-methyl-hex-4-enoic acid

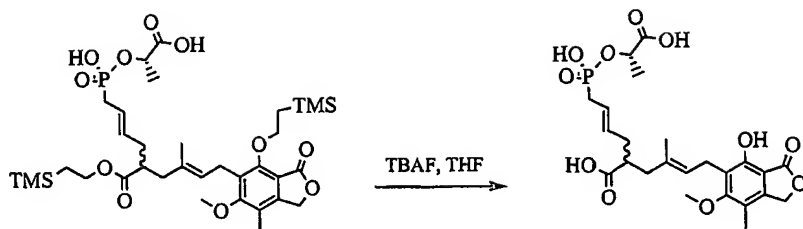
A solution of 2-{4-[(1-ethoxycarbonyl-ethoxy)-methoxy-phosphoryl]-but-2-enyl}-6-[6-methoxy-7-methyl-3-oxo-4-(2-trimethylsilyl-ethoxy)-1,3-dihydro-isobenzofuran-5-yl]-4-methyl-hex-4-enoic acid 2-trimethylsilyl-ethyl ester (14 mg, 0.018 mmol) in THF (1 mL) was stirred with a 1M solution of TBAF in THF (55 µL, 0.055 mmol) for 1 hour. The reaction mixture was concentrated, acidified with 1N HCl and extracted with EtOAc. The organic layer was washed with brine and dried. The product was purified by silica gel column chromatography EtOH-EtOAc (0-10%). Further purification was performed by dissolving the product in CH₂Cl₂ and passing the compound through a 13 mm Acrodisc syringe filter with a 0.45 µm Nylon membrane to

provide 8 mg (77%) of the product. ^1H NMR (300 MHz, CDCl_3) δ 0.92 (t, 3H, $J=7$ Hz), 1.30 (d, 3H, $J=8$ Hz), 1.79 (s, 3H), 2.10-2.39 (m, 4H), 2.15 (s, 3H), 2.53 (d, 1H, $J=8$ Hz), 2.65 (d, 1H, $J=22$ Hz), 2.68 (d, 1H, $J=22$ Hz), 3.38 (d, 2H, $J=7$ Hz), 3.70 and 3.74 (d, 3H, $J=11$ Hz), 3.76 (s, 3H), 4.07 (m, 2H), 4.96 (dq, 1H, $J=7$ Hz), 5.20 (s, 2H), 5.27 (br t, 1H, $J=7$ Hz), 5.33-5.55 (m, 2H), 7.51-7.56 (m, 1H), 7.68-7.74 (m, 1H) ppm; ^{31}P (121.4 MHz, CDCl_3) δ 29.0, 30.1 ppm; MS (m/z) 569.2 $[\text{M}+\text{H}]^+$, 591.3 $[\text{M}+\text{Na}]^+$.



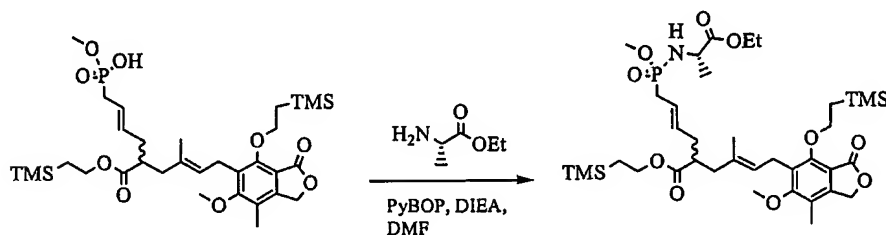
2-{4-[(1-Carboxy-ethoxy)-hydroxy-phosphoryl]-but-2-enyl}-6-[6-methoxy-7-methyl-3-oxo-4-(2-trimethylsilyl-ethoxy)-1,3-dihydro-isobenzofuran-5-yl]-4-methyl-hex-4-enoic acid 2-trimethylsilyl-ethyl ester

A solution of 2-{4-[(1-ethoxycarbonyl-ethoxy)-methoxy-phosphoryl]-but-2-enyl}-6-[6-methoxy-7-methyl-3-oxo-4-(2-trimethylsilyl-ethoxy)-1,3-dihydro-isobenzofuran-5-yl]-4-methyl-hex-4-enoic acid 2-trimethylsilyl-ethyl ester (12 mg, 0.016 mmol) in *tert*-butylamine (1 mL, 9.6 mmol) was heated at 65°C for 16 hours. The solution was allowed to cool to room temperature and concentrated to provide the crude product as an oil. ^1H NMR (300 MHz, CDCl_3) δ 0.03 (s, 9H), 0.04 (s, 9H), 0.86-0.98 (m, 2H), 1.22-1.33 (m, 2H), 1.50 (d, 3H, $J=7$ Hz), 1.78 (s, 3H), 2.05-2.30 (m, 4H), 2.10 (s, 3H), 2.48-2.63 (m, 3H), 3.40 (d, 2H, $J=7$ Hz), 3.76 (s, 3H), 4.08 (appt t, 2H, $J=9$ Hz), 4.25-4.33 (m, 2H), 4.75-4.84 (m, 1H), 5.13 (s, 2H), 5.15-5.23 (m, 1H), 5.33-5.55 (m, 2H) ppm; ^{31}P (121.4 MHz, CDCl_3) δ 28.9 ppm; MS (m/z) 725.3 $[\text{M}-\text{H}]^-$.



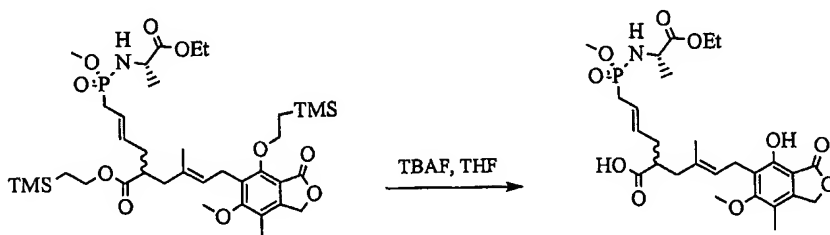
2-{4-[(1-Carboxy-ethoxy)-hydroxy-phosphoryl]-but-2-enyl}-6-(4-hydroxy-6-methoxy-7-methyl-3-oxo-1,3-dihydro-isobenzofuran-5-yl)-4-methyl-hex-4-enoic acid

A solution of crude 2-{4-[(1-carboxy-ethoxy)-hydroxy-phosphoryl]-but-2-enyl}-6-[6-methoxy-7-methyl-3-oxo-4-(2-trimethylsilylanyl-ethoxy)-1,3-dihydro-isobenzofuran-5-yl]-4-methyl-hex-4-enoic acid 2-trimethylsilylanyl-ethyl ester (AC-2101-59) and tetrabutylammonium fluoride in THF (1M, 54 μ L, 0.054 mmol) was stirred with THF (1 mL) for 2 hours at ambient temperature, when more tetrabutylammonium fluoride in THF (54 μ L, 0.054 mmol) was added. The reaction was stirred for an additional 16 hours, by which time the reaction was complete. The reaction mixture was concentrated *in vacuo* and the product was purified by RP HPLC using a Phenomenex Synergi 5 μ Hydro RP 80A column (50 x 21.2 mm) with eluents of H₂O, 0.1% TFA-CH₃CN, 0.1% TFA to provide the product (8.0 mg) as a clear oil. ¹H NMR (300 MHz, CDCl₃) δ 1.51 (d, 3H, *J* = 7 Hz), 1.79 (s, 3H), 2.05- 2.40 (m, 4H), 2.11 (s, 3H), 2.49- 2.71 (m, 3H), 3.38 (d, 2H, *J* = 6 Hz), 3.76 (s, 3H), 4.85 (br s, 1H), 5.20 (s, 2H), 5.21- 5.30 (m, 1H), 5.33- 5.63 (m, 2H) ppm; ³¹P (121.4 MHz, CDCl₃) δ 27.7 ppm; MS (*m/z*) 525.2 [M-H]⁻.



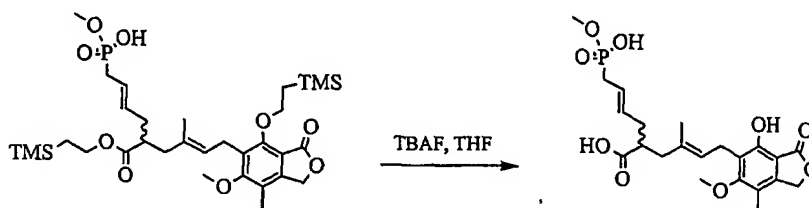
2-{4-[(1-Ethoxycarbonyl-ethylamine)-methoxy-phosphoryl]-but-2-enyl}-6-[6-methoxy-7-methyl-3-oxo-4-(2-trimethylsilanyl-ethoxy)-1,3-dihydro-isobenzofuran-5-yl]-4-methyl-hex-4-enoic acid 2-trimethylsilanyl-ethyl ester

A solution of 2-[4-(hydroxy-methoxy-phosphoryl)-but-2-enyl]-6-[6-methoxy-7-methyl-3-oxo-4-(2-trimethylsilanyl-ethoxy)-1,3-dihydro-isobenzofuran-5-yl]-4-methyl-hex-4-enoic acid 2-trimethylsilanyl-ethyl ester (20 mg, 0.030 mmol), PyBOP (62.4 mg, 0.120 mmol) in DMF (1.0 mL) was stirred with L-alanine ethyl ester hydrochloride (18 mg, 0.12 mmol) and DIEA (26 μ L, 0.15 mmol) at ambient temperature for 1 hour, when complete consumption of the starting materials was observed. The reaction was worked up by addition of water until the reaction solution became cloudy. DMF was added dropwise until the mixture became clear again. The reaction mixture was filtered through Acrodisc (13 mm syringe filter with a 0.45 μ m Nylon membrane) and purified by RP HPLC using a Phenomenex Synergi 5 μ Hydro RP 80A column (50 x 21.2 mm), eluting with water and acetonitrile. The fractions containing the product were pooled together and concentrated *in vacuo* to remove the acetonitrile. The remaining solution was saturated with sodium chloride and extracted with EtOAc and acetonitrile to provide 7.2 mg of the product. ^1H NMR (300 MHz, CDCl_3) δ 0.03 (s, 9H), 0.05 (s, 9H), 0.923 (appt t, 2H, $J=8$ Hz), 1.18- 1.31 (m, 5H), 1.41 (t, 3H, $J=7$ Hz), 1.78 (s, 3H), 2.03- 2.36 (m, 4H), 2.18 (s, 3H), 2.43- 2.63 (m, 3H), 3.10- 3.30 (m, 1H), 3.40 (d, 2H, $J=7$ Hz), 3.62 and 3.65 (d, 3H, $J=11$ Hz), 3.76 (s, 3H), 4.03-4.12 (m, 2H), 4.20 (dq, 2H, $J=2, 7$ Hz), 4.29 (appt t, 2H, $J=8$ Hz), 5.12 (s, 2H), 5.18- 5.28 (m, 1H), 5.33- 5.67 (m, 2H) ppm; ^{31}P (121.4 MHz, CDCl_3) δ 30.4, 31.2 ppm; MS (m/z) 790.4 $[\text{M}+\text{Na}]^+$.



2-{4-[(1-Ethoxycarbonyl-ethylamine)-methoxy-phosphoryl]-but-2-enyl}-6-(4-hydroxy-6-methoxy-7-methyl-3-oxo-1,3-dihydro-isobenzofuran-5-yl)-4-methyl-hex-4-enoic acid

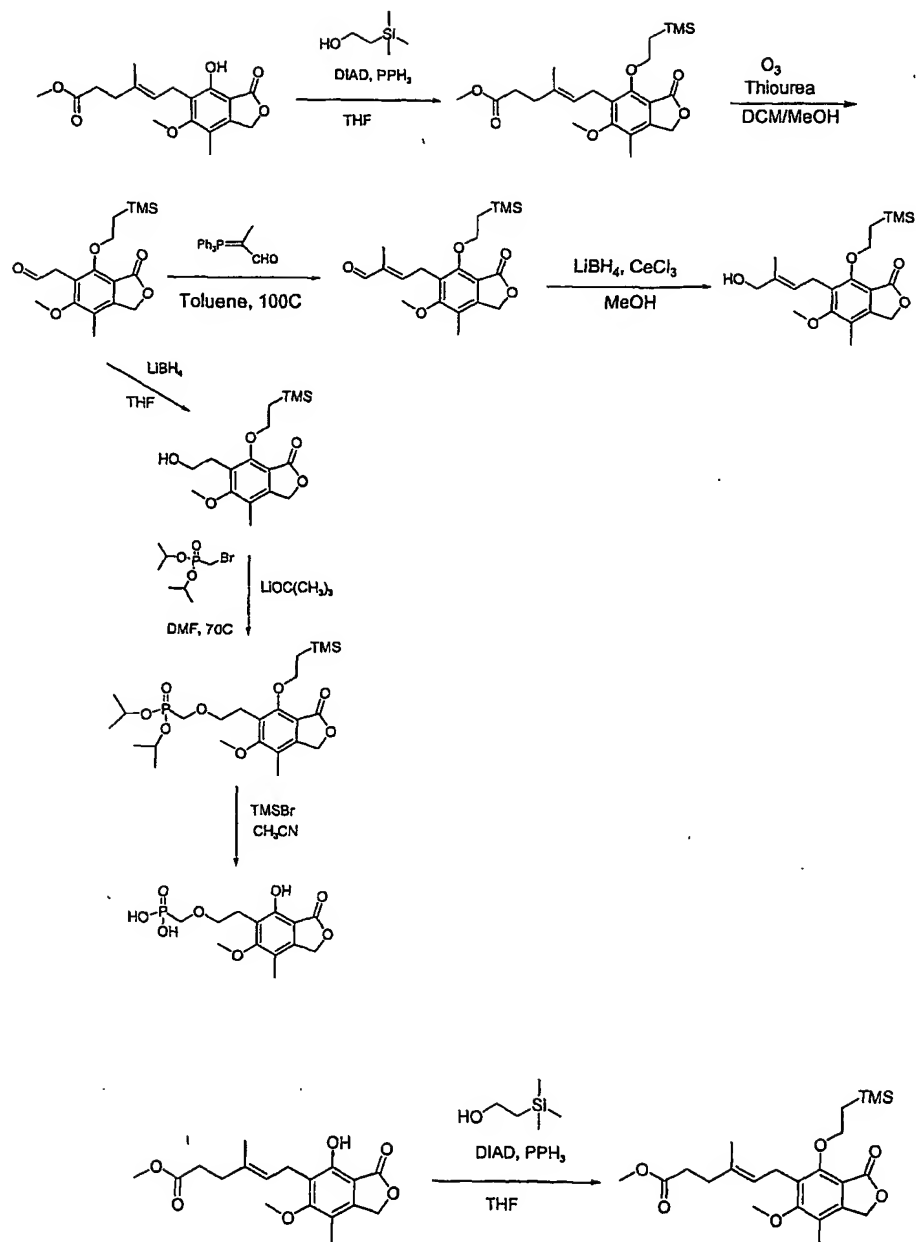
To a solution of 2-{4-[(1-ethoxycarbonyl-ethylamine)-methoxy-phosphoryl]-but-2-enyl}-6-[6-methoxy-7-methyl-3-oxo-4-(2-trimethylsilyl-ethoxy)-1,3-dihydro-isobenzofuran-5-yl]-4-methyl-hex-4-enoic acid 2-trimethylsilyl-ethyl ester (7.2 mg, 9.38 mmol) in THF (1 mL) was added TBAF (40 μ L, 1M solution in THF) at room temperature. The reaction mixture was stirred for 20 minutes, when the starting material was completely converted to the desired product as judged by LCMS. The reaction mixture was dried *in vacuo* and re-dissolved in DMF. The product was purified by RP HPLC using a Phenomenex Synergi 5 μ Hydro RP 80A column (50 x 21.2 mm) with eluents of H₂O-CH₃CN. The fractions containing the desired product were pooled and further purified on Dowex 50WX8-400 packed on a 4.5 cm x 2 cm column to elute the sodium salt at H₂O- MeOH (1:1), providing 3.2 mg of the desired product. ¹H NMR (300 MHz, CD₃OD) δ 1.26 (dd, 3H, *J* = 4, 7 Hz), 1.37 (t, 3H, *J* = 8 Hz), 1.80 (s, 3H), 2.00- 2.22 (m, 4H), 2.10 (s, 3H), 2.25- 2.60 (m, 3H), 3.37 (d, 2H, *J* = 7 Hz), 3.60 and 3.65 (d, 3H, *J* = 11 Hz), 3.74 (s, 3H), 3.83- 3.96 (m, 1H), 4.18 (q, 2H, *J* = 8 Hz), 5.15 (s, 2H), 5.25- 5.42 (m, 2H), 5.55- 5.69 (m, 1H) ppm; ³¹P (121.4 MHz, CD₃OD) δ 33.8, 34.2 ppm; MS (*m/z*) 568.2 [M+H]⁺, 590.3 [M+Na]⁺.



6-(4-Hydroxy-6-methoxy-7-methyl-3-oxo-1,3-dihydro-isobenzofuran-5-yl)-2-[4-(hydroxy-methoxy-phosphoryl)-but-2-enyl]-4-methyl-hex-4-enoic acid

To a solution of 2-[4-(hydroxy-methoxy-phosphoryl)-but-2-enyl]-6-[6-methoxy-7-methyl-3-oxo-4-(2-trimethylsilyl-ethyl-ethoxy)-1,3-dihydro-isobenzofuran-5-yl]-4-methyl-hex-4-enoic acid 2-trimethylsilyl-ethyl ester (11

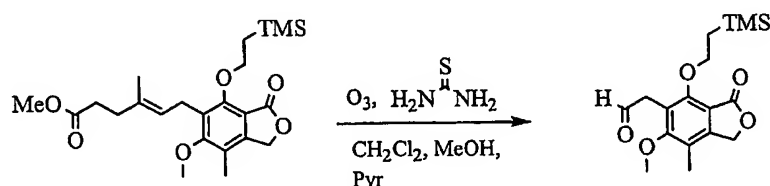
mg, 0.016 mmol) in THF (1 mL) was added TBAF (50 μ L, 1M solution in THF) at room temperature. The solution was stirred for 16 hours and concentrated. The solution was dried under reduced pressure and re-suspended in DMF (0.8 mL) and water (0.25 mL). The solution was filtered through Acrodisc (13 mm syringe filter with a 0.45 μ m Nylon membrane) and purified by RP HPLC using a Phenomenex Synergi 5 μ Hydro RP 80A column (50 x 21.2 mm) with eluents of H₂O, 0.1% TFA-CH₃CN, 0.1% TFA. The product from the column was subjected to ion exchange chromatography (Sodium salt form of Dowex 50WX8-400) using a 2 x 4.5 cm column eluting with H₂O-MeOH (1:1) to provide 7.5 mg of the desired product as an oil. ¹H NMR (300 MHz, CDCl₃) δ 1.80 (s, 3H), 2.01- 2.29 (m, 5H), 2.11 (s, 3H), 2.35 (d, 2H, *J*= 22 Hz), 3.38 (d, 2H, *J*= 7 Hz), 3.53 (d, 3H, *J*= 11 Hz), 3.75 (s, 3H), 5.19 (s, 2H), 5.26 (t, 1H, *J*= 6 Hz), 5.43- 5.54 (m, 2H) ppm; ³¹P (121.4 MHz, CDCl₃) δ 23.5 ppm; MS (*m/z*) 469.2 [M+H]⁺, 491.3 [M+Na]⁺.



6-[6-Methoxy-7-methyl-3-oxo-4-(2-trimethylsilyl-ethoxy)-1,3-dihydro-isobenzofuran-5-yl]-4-methyl-hex-4-enoic acid methyl ester

To a solution of 6-(4-hydroxy-6-methoxy-7-methyl-3-oxo-1,3-dihydro-isobenzofuran-5-yl)-4-methyl-hex-4-enoic acid methyl ester (222 mg, 0.66 mmol), triphenylphosphine (260 mg, 0.996 mmol), and diethyl azodicarboxylate (173 mg, 0.996 mmol) in THF (3 mL) at 0°C was added a solution of 2-trimethylsilyl-ethanol (142 μL, 0.996 mmol) in THF (3 mL). The resulting yellow

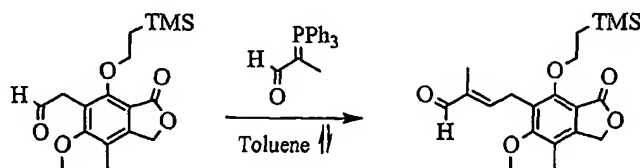
solution was allowed to warm to room temperature and stirred overnight. The reaction was concentrated to dryness and ether and hexanes were added. Triphenylphosphine oxide was removed by filtration and the filtrate was concentrated and purified by silica gel chromatography to provide 248 mg of the desired product as a colorless oil. ^1H NMR (300 MHz, CDCl_3) δ 0.03 (s, 9H), 1.18- 1.30 (m, 2H), 1.81 (s, 3H), 2.18 (s, 3H), 2.25- 2.33 (m, 2H), 2.37- 2.45 (m, 2H), 3.42 (d, 2H, $J=7$ Hz), 3.62 (s, 3H), 3.77 (s, 3H), 4.25- 4.35 (m, 2H), 5.13 (s, 2H), 5.12- 5.22 (m, 1H) ppm.



[6-Methoxy-7-methyl-3-oxo-4-(2-trimethylsilyl-ethoxy)-1,3-dihydro-isobenzofuran-5-yl]-acetaldehyde

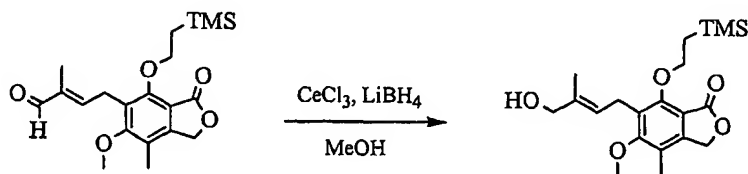
A solution of 6-[6-methoxy-7-methyl-3-oxo-4-(2-trimethylsilyl-ethoxy)-1,3-dihydro-isobenzofuran-5-yl]-4-methyl-hex-4-enoic acid methyl ester (618 mg, 1.42 mmol) in MeOH (10 mL), CH_2Cl_2 (10 mL) and pyridine (50 μL , 0.618 mmol) was cooled to -70°C using a dry ice/ acetone bath according to the procedure of Smith, D. B. *et al.*, *J. Org. Chem.*, 1996, 61, 6, 2236. A stream of ozone was bubbled through the reaction *via* a gas dispersion tube until the reaction became blue in color (15 minutes). The ozone line was replaced with a stream of nitrogen and bubbling continued for another 15 minutes, by which time the blue color had disappeared. To this solution, thiourea (75.7 mg, 0.994 mmol) was added in one portion at -70°C , and the cooling bath was removed. The reaction was allowed to warm to room temperature and stirred for 15 hours. The reaction was worked up by filtration to remove solid thiourea S-dioxide, and then partitioned between CH_2Cl_2 and water. The organic layer was removed. The aqueous layer was washed with CH_2Cl_2 one more time, and the organic extracts were combined. The organic layer was washed with aqueous 1N HCl, saturated NaHCO_3 and brine. The organic extracts were dried *in vacuo* and the residue was purified to by silica gel chromatography to afford 357 mg (75 %) of the

product as a white solid. ^1H NMR (300 MHz, CDCl_3) δ -0.01 (s, 9H), 1.05- 1.15 (m, 2H), 2.15 (s, 3H), 3.69 (s, 3H), 3.78 (d, 2H, J = 1 Hz), 4.27- 4.39 (m, 2H), 5.11 (s, 2H), 9.72 (d, 1H, J = 1 Hz) ppm.



4-[6-Methoxy-7-methyl-3-oxo-4-(2-trimethylsilyl-ethoxy)-1,3-dihydro-isobenzofuran-5-yl]-2-methyl-but-2-enal

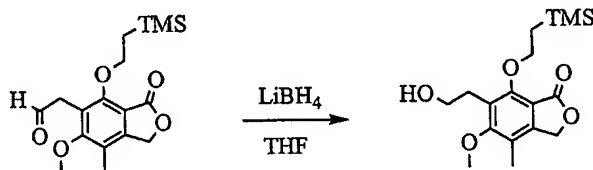
[6-Methoxy-7-methyl-3-oxo-4-(2-trimethylsilyl-ethoxy)-1,3-dihydro-isobenzofuran-5-yl]-acetaldehyde (70 mg, 0.21 mmol) in toluene (2 mL) was heated at 100 °C with 2-(triphenyl-phosphanylidene)-propionaldehyde (72.9 mg, 0.23 mmol) overnight. A second portion of 2-(triphenyl-phosphanylidene)-propionaldehyde (33 mg, 0.11 mmol) was added and the reaction mixture was heated for an additional day. After concentration, the residue was purified by silica gel chromatography to provide 54 mg (83%) of the desired product as a pale yellow oil. ^1H NMR (300 MHz, CDCl_3) δ 0.00 (s, 9H), 1.10- 1.21 (m, 2H), 1.87 (s, 3H), 2.16 (s, 3H), 3.67- 3.76 (m, 2H), 3.74 (s, 3H), 4.27- 4.39 (m, 2H), 5.11 (s, 2H), 6.40- 6.48 (m, 1H), 9.2 (s, 1H) ppm.



6-(4-Hydroxy-3-methyl-but-2-enyl)-5-methoxy-4-methyl-7-(2-trimethylsilyl-ethoxy)-3H-isobenzofuran-1-one

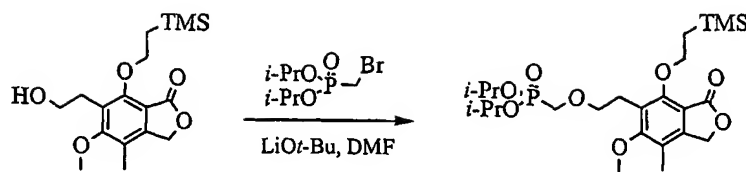
A solution of 4-[6-methoxy-7-methyl-3-oxo-4-(2-trimethylsilyl-ethoxy)-1,3-dihydro-isobenzofuran-5-yl]-2-methyl-but-2-enal (103 mg, 0.27 mmol) in methanol (5 mL) was cooled to 0°C. A solution of CeCl_3 (0.68 mL, MeOH: H_2O , 9:1) was added, followed by LiBH_4 (0.14 mL, 0.28 mmol of a 2M

solution in THF). The ice bath was removed and the reaction mixture was allowed to warm to room temperature. The reaction mixture was stirred for an additional 40 minutes whereupon TLC indicated complete consumption of starting aldehyde. The reaction was worked up by addition of aqueous 1N HCl (0.5 mL) and the product was extracted with CH₂Cl₂. The organic layer was washed with saturated aqueous sodium bicarbonate solution and brine. The organic layer was concentrated under reduced pressure and the residue was purified by silica gel chromatography to provide 100 mg (97%) of the product as a clear liquid. ¹H NMR (300 MHz, CDCl₃) δ 0.00 (s, 9H), 1.20 (dd, 2H, *J*= 7, 8 Hz), 1.81 (s, 3H), 2.13 (s, 3H), 3.38- 3.50 (m, 2H), 3.74 (s, 3H), 3.95 (s, 2H), 4.27 (dd, 2H, *J*= 7, 8 Hz), 5.08 (s, 2H), 5.17- 5.44 (m, 1H) ppm.



6-(2-Hydroxy-ethyl)-5-methoxy-4-methyl-7-(2-trimethylsilyl-ethoxy)-3H-isobenzofuran-1-one

To a solution of [6-methoxy-7-methyl-3-oxo-4-(2-trimethylsilyl-ethoxy)-1,3-dihydro-isobenzofuran-5-yl]-acetaldehyde (97 mg, 0.29 mmol) in THF (5 mL) was added an aliquot of a 2 M LiBH₄ in THF (150 μL, 0.300 mmol). The reaction mixture was stirred at room temperature for 1 hour when complete consumption of the starting materials was observed by TLC. The reaction mixture was worked up by addition of an aqueous 1N HCl solution and extraction with EtOAc. The organic layer was dried *in vacuo* and the residue was purified by silica gel chromatography to provide the product. ¹H NMR (300 MHz, CDCl₃) δ 0.00 (s, 9H), 1.20 (dd, 2H, *J*=7, 9 Hz), 2.07 (br s, 1H), 2.14 (s, 3H), 2.97 (t, 2H, *J*= 6 Hz), 3.76 (t, 2H, *J*= 6 Hz), 3.77 (s, 3H), 4.32 (dd, 2H, *J*= 7, 8 Hz), 5.08 (s, 2H) ppm.

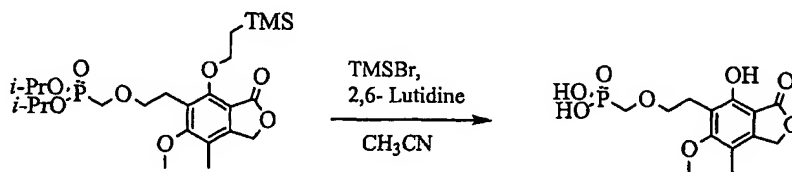


{2-[6-Methoxy-7-methyl-3-oxo-4-(2-trimethylsilyl-ethoxy)-1,3-dihydro-isobenzofuran-5-yl]-ethoxymethyl}-phosphonic acid diisopropyl ester

A mixture of 6-(2-hydroxy-ethyl)-5-methoxy-4-methyl-7-(2-trimethylsilyl-ethoxy)-3H-isobenzofuran-1-one (79 mg, 0.23 mmol) was heated with bromomethylphosphonic acid diisopropyl ester (120 mg, 0.46 mmol) in the presence of lithium *t*-butoxide (22 mg, 0.27 mmol) in DMF (2 mL) at 70 °C overnight. The reaction mixture was purified by RP HPLC (acetonitrile and 0.1% aqueous CF₃COOH) to provide the desired product. ¹H NMR (300 MHz, CDCl₃) δ 0.00 (s, 9H), 1.13- 1.25 (m, 2H), 1.26 (t, 12H, *J* = 6 Hz), 2.12 (s, 3H), 2.98 (t, 2H, *J* = 7 Hz), 3.60- 3.73 (m, 4H), 3.77 (s, 3H), 4.05- 4.16 (m, 2H), 4.62- 4.74 (m, 2H), 5.07 (s, 2H) ppm; MS (*m/z*) 539 [M+Na]⁺.

Example 296

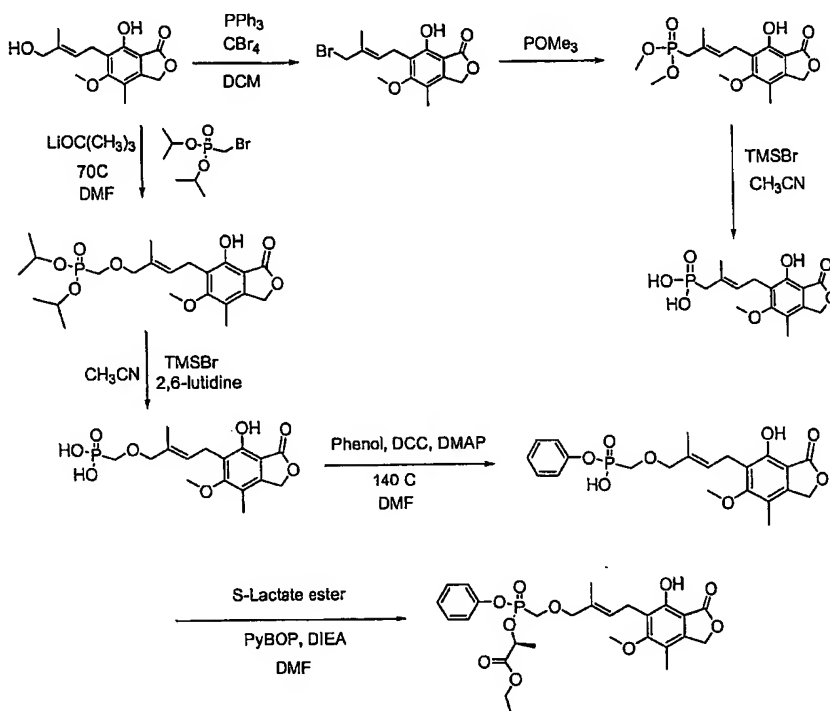
Representative compounds of the invention can be prepared as illustrated below.



[2-(4-Hydroxy-6-methoxy-7-methyl-3-oxo-1,3-dihydro-isobenzofuran-5-yl)-ethoxymethyl]-phosphonic acid

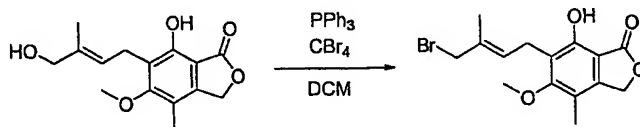
To a solution of {2-[6-methoxy-7-methyl-3-oxo-4-(2-trimethylsilyl-ethoxy)-1,3-dihydro-isobenzofuran-5-yl]-ethoxymethyl}-phosphonic acid diisopropyl ester (7.5 mg, 0.014 mmol) in acetonitrile (2 mL) and 2,6-lutidine (25 μL, 0.21 mmol) was added trimethylsilyl bromide (27 μL, 0.21 mmol) at room temperature. The reaction was allowed to proceed for 18 hours when completion of the reaction was indicated by LCMS. The reaction was quenched

by addition of MeOH and concentration. The residue was purified by RP-HPLC using a C18 column. The collected product was dissolved in a solution of 10% TFA/ CH₂Cl₂ to assure complete deprotection. The reaction mixture was lyophilized to provide the desired product. ¹H NMR (300 MHz, CD₃OD) δ 2.12 (s, 3H), 2.98 (t, 2H, *J*= 7 Hz), 3.66- 3.76 (m, 4H), 3.78 (s, 3H), 5.21 (s, 2H) ppm; MS (*m/z*) 331 [M-H].



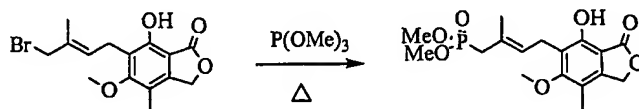
Example 297

Representative compounds of the invention can be prepared as illustrated below.



6-(4-Bromo-3-methyl-but-2-enyl)-7-hydroxy-5-methoxy-4-methyl-3H-isobenzofuran-1-one

Polymer-supported triphenylphosphine (3 mmol/g, 0.5 g) was soaked in dichloromethane (10mL) for 1 hour 7-Hydroxy-6-(4-hydroxy-3-methyl-but-2-enyl)-5-methoxy-4-methyl-3H-isobenzofuran-1-one (100 mg, 0.36 mmol) and carbon tetrabromide (143 mg, 0.43 mmol) were added sequentially and the mixture was shaken for 1 hour at room temperature. More carbon tetrabromide (143 mg, 0.43 mmol) was added and the mixture was shaken further for 1 hour. The mixture was filtered and the filtrate was concentrated. The residue was chromatographed on silica gel (0 % to 60 % ethyl acetate / hexanes) to afford 6-(4-bromo-3-methyl-but-2-enyl)-7-hydroxy-5-methoxy-4-methyl-3H-isobenzofuran-1-one as an oil (52 mg, 42 %); ^1H NMR (300 MHz, CDCl_3) δ 1.95 (s, 3H), 2.16 (s, 3H), 3.44 (d, $J = 7.2$, 2H), 3.78 (s, 3H), 3.98 (s, 2H), 5.21 (s, 2H), 5.68 (t, $J = 7.2$ Hz, 1H), 7.71 (brs, 1H) ppm.

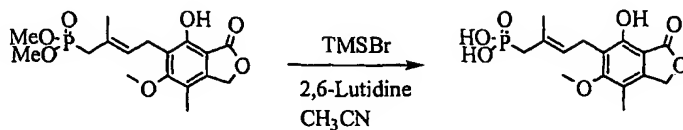


[4-(4-Hydroxy-6-methoxy-7-methyl-3-oxo-1,3-dihydro-isobenzofuran-5-yl)-2-methyl-but-2-enyl]-phosphonic acid dimethyl ester

A solution of 6-(4-bromo-3-methyl-but-2-enyl)-7-hydroxy-5-methoxy-4-methyl-3H-isobenzofuran-1-one (33 mg, 0.097 mmol) in trimethylphosphite (1.0 mL, 8.5 mmol) was heated to 100 °C for 1 hour, whereupon complete reaction was indicated by LCMS. The reaction was worked up by removal of the excess reagent under reduced pressure and the residue was purified by silica gel chromatography using EtOAc-hexanes (20-100%) to provide 20 mg (60%) of the desired product. ^1H NMR (300 MHz, CDCl_3) δ 1.90 (s, 3H), 2.09 (s, 3H), 2.48 (d, 2H, $J = 22$ Hz), 3.38 (t, 2H, $J = 6$ Hz), 3.64 (d, 6H, $J = 11$ Hz), 3.72 (s, 3H), 5.14 (s, 2H), 5.33 (q, 1H, $J = 6$ Hz), 7.65 (br s, 1H) ppm; MS (m/z) 371 $[\text{M}+\text{H}]^+$.

Example 298

Representative compounds of the invention can be prepared as illustrated below.

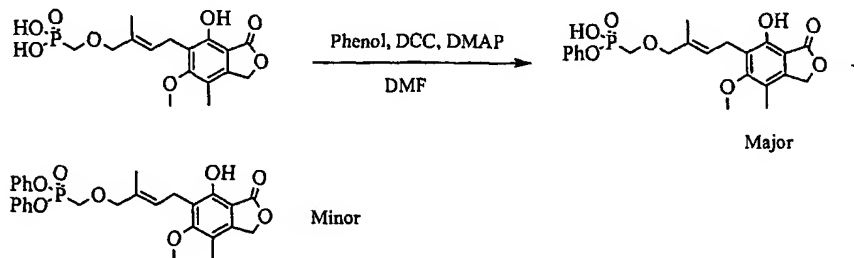


[4-(4-Hydroxy-6-methoxy-7-methyl-3-oxo-1,3-dihydro-isobenzofuran-5-yl)-2-methyl-but-2-enyl]-phosphonic acid

To a solution of [4-(4-hydroxy-6-methoxy-7-methyl-3-oxo-1,3-dihydro-isobenzofuran-5-yl)-2-methyl-but-2-enyl]-phosphonic acid dimethyl ester (18 mg, 0.049 mmol) in acetonitrile (2 mL) was added TMSBr (63 μ L, 0.49 mmol) and 2,6-lutidine (85 μ L, 0.73 mmol) at 0°C. The reaction solution was allowed to warm to room temperature and stirred for 2 hours when completion of the reaction was observed by LCMS. The reaction was cooled to 0°C and quenched by the addition of MeOH. The reaction mixture was concentrated under reduced pressure and the residue was purified by RP HPLC using a C18 column with a gradient of H₂O-acetonitrile (5-0%) over 20 minutes to provide 12.2 mg (73%) of the product. ¹H NMR (300 MHz, CD₃OD) δ 1.95 (s, 3H), 2.15 (s, 3H), 2.48 (d, 2H, J = 22 Hz), 3.44 (t, 2H, J = 6 Hz), 3.79 (s, 3H), 5.24 (s, 2H), 5.38 (q, 1H, J = 7 Hz), 6.87 (br s, 1H) ppm; MS (m/z) 341 [M-H]⁻.

Example 299

Representative compounds of the invention can be prepared as illustrated below.

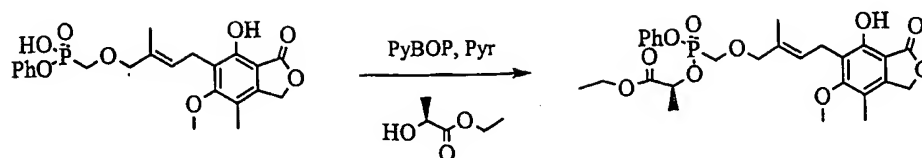


[4-(4-Hydroxy-6-methoxy-7-methyl-3-oxo-1,3-dihydro-isobenzofuran-5-yl)-2-methyl-but-2-enyloxymethyl]-phosphonic acid monophenyl ester and [4-(4-Hydroxy-6-methoxy-7-methyl-3-oxo-1,3-dihydro-isobenzofuran-5-yl)-2-methyl-but-2-enyloxymethyl]-phosphonic acid diphenyl ester

To a solution of [4-(4-hydroxy-6-methoxy-7-methyl-3-oxo-1,3-dihydro-isobenzofuran-5-yl)-2-methyl-but-2-enyloxymethyl]-phosphonic acid (49 mg, 0.13 mmol) in DMF (0.4 mL) and phenol (62 mg, 0.65 mmol) was added dicyclohexyl carbodiimide (107 mg, 0.52 mmol) and DMAP (8 mg, 0.065 mmol) in DMF (0.6 mL), slowly at 0°C. The reaction was allowed to warm to room temperature and heated to 140°C for 10 hours. After cooling to room temperature the mixture was filtered and extracted with aqueous 1N NaOH solution. The aqueous layer was acidified with aqueous 1N HCl and extracted with EtOAc. The organic layer was dried over Na₂SO₄ and concentrated to dryness. The residue was purified by RP HPLC to provide 18.5 mg of [4-(4-hydroxy-6-methoxy-7-methyl-3-oxo-1,3-dihydro-isobenzofuran-5-yl)-2-methyl-but-2-enyloxymethyl]-phosphonic acid monophenyl ester, as a pale yellow solid and 4.1 mg of [4-(4-hydroxy-6-methoxy-7-methyl-3-oxo-1,3-dihydro-isobenzofuran-5-yl)-2-methyl-but-2-enyloxymethyl]-phosphonic acid diphenyl ester (minor product) also as a pale yellow solid. Major product: ¹H NMR (300 MHz, CD₃OD) δ 1.82 (s, 3H), 2.16 (s, 3H), 3.46 (d, 2H, *J* = 7 Hz), 3.70 (d, 2H, *J* = 8 Hz), 3.77 (s, 3H), 3.96 (s, 2H), 5.25 (s, 2H), 5.52 (t, 1H, *J* = 8 Hz), 7.10-7.21 (m, 3H), 7.30 (t, 2H, *J* = 8 Hz) ppm; ³¹P (121.4 MHz, CD₃OD) δ 17.3 ppm; MS (*m/z*) 449.0 [M+H]⁺, 471.2 [M+Na]⁺. Minor product: ¹H NMR (300 MHz, CD₃OD) δ 1.82 (s, 3H), 2.15 (s, 3H), 3.47 (d, 2H, *J* = 7 Hz), 3.77 (s, 3H), 3.98-4.06 (m, 4H), 5.25 (s, 2H), 5.50-5.61 (m, 1H), 7.10-7.25 (m, 6H), 7.30-7.41 (m, 4H) ppm; ³¹P (121.4 MHz, CD₃OD) δ 16.3 ppm; MS (*m/z*) 525.2 [M+H]⁺, 547.2 [M+Na]⁺.

Example 300

Representative compounds of the invention can be prepared as illustrated below.

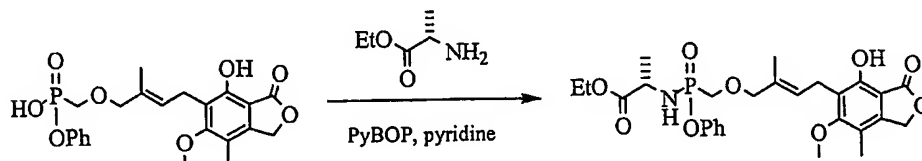


2-[[4-(4-Hydroxy-6-methoxy-7-methyl-3-oxo-1,3-dihydro-isobenzofuran-5-yl)-2-methyl-but-2-enyloxymethyl]-phenoxy-phosphinoyloxy}-propionic acid ethyl ester

To a solution of [4-(4-hydroxy-6-methoxy-7-methyl-3-oxo-1,3-dihydro-isobenzofuran-5-yl)-2-methyl-but-2-enyloxymethyl]-phosphonic acid monophenyl ester (18.5 mg, 0.040 mmol) and ethyl (S)-(-)-lactate (47 μ L, 0.400 mmol) in pyridine (0.5 mL) was added PyBOP (32 mg, 0.060 mmol). The solution was stirred at room temperature for 1 hour, when an additional portion of PyBOP (21 mg, 0.040 mmol) was added. The solution was stirred for another hour and concentrated. The residue was purified by HPLC to provide 7.5 mg of the desired product as a clear oil. ^1H NMR (300 MHz, CD_3OD) δ 1.22 and 1.25 (t, 3H, $J = 7$ Hz), 1.42 and 1.50 (d, 3H, $J = 7$ Hz), 1.82 and 1.83 (s, 3H), 2.16 (s, 3H), 3.47 (d, 2H, $J = 7$ Hz), 3.78 (s, 3H), 3.89 (d, 1H, $J = 8$ Hz), 3.93-4.02 (m, 3H), 4.10-4.22 (m, 2H), 4.94-5.08 (m, 1H), 5.25 (s, 2H), 5.50-5.60 (m, 1H), 7.15-7.27 (m, 3H), 7.33-7.41 (m, 2H) ppm; ^{31}P (121.4 MHz, CD_3OD) δ 18.9, 20.3 ppm (diastereomers at phosphorus); MS (m/z) 549.2 $[\text{M}+\text{H}]^+$, 571.3 $[\text{M}+\text{Na}]^+$.

Example 301

Representative compounds of the invention can be prepared as illustrated below.

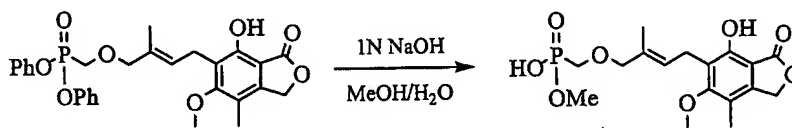


2-{{4-(4-Hydroxy-6-methoxy-7-methyl-3-oxo-1,3-dihydro-isobenzofuran-5-yl)-2-methyl-but-2-enyloxymethyl]-phenoxy-phosphinoylamino}-propionic acid ethyl ester

To a solution of [4-(4-hydroxy-6-methoxy-7-methyl-3-oxo-1,3-dihydro-isobenzofuran-5-yl)-2-methyl-but-2-enyloxymethyl]-phosphonic acid monophenyl ester (20 mg, 0.045 mmol) and L-alanine ethyl ester hydrochloride (68.5mg, 0.45mmol) in pyridine (1.0 mL) was added PyBOP (70mg, 0.14mmol). After stirring overnight, the mixture was concentrated and the residue purified by RP HPLC using a C18 column with a gradient of H₂O, 0.1% TFA-acetonitrile, 0.1% TFA to provide 3.6 mg of the product as a colorless gel. ¹H NMR (300 MHz, CD₃OD) δ 1.17-1.3 (m, 6H), 1.8-1.9 (m, 3H), 2.16 (s, 3H), 3.17 (m, 1H), 3.47 (d, 2H), 3.72-3.8 (m, 5H), 3.92-4.2 (m, 4H), 5.25 (s, 2H), 5.54 (m, 1H), 7.18 (m, 3H), 7.33 (m, 2H) ppm; ³¹P (121.4 MHz, CD₃OD) δ 24.1, 25.0 ppm (diastereomers at phosphorus); MS (*m/z*) 546.2 [M-H]⁺.

Example 302

Representative compounds of the invention can be prepared as illustrated below.



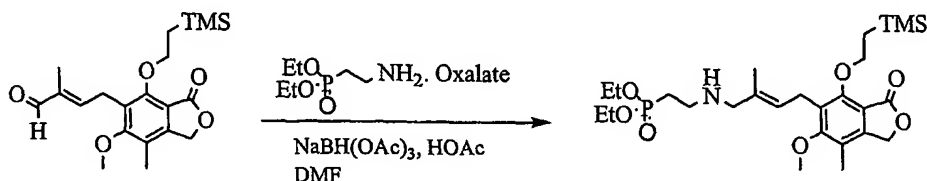
[4-(4-Hydroxy-6-methoxy-7-methyl-3-oxo-1,3-dihydro-isobenzofuran-5-yl)-2-methyl-but-2-enyloxymethyl]-phosphonic acid monomethyl ester

To a solution of [4-(4-hydroxy-6-methoxy-7-methyl-3-oxo-1,3-dihydro-isobenzofuran-5-yl)-2-methyl-but-2-enyloxymethyl]-phosphonic acid diphenyl ester (53 mg, 0.1 mmol) in methanol (0.5 mL) was added an aqueous solution of 1N NaOH (300 µL). After stirring overnight, the mixture was concentrated and the residue purified by RP HPLC using a C18 column with a gradient of H₂O, 0.1% TFA-acetonitrile, 0.1% TFA to provide 5 mg of the product as a colorless gel, together with the phosphonic acid monophenyl ester (7mg) and the phosphonic acid dimethyl ester (14.5mg). ¹H NMR (300 MHz, CD₃OD) δ 1.84

(s, 3H), 2.16 (s, 3H), 3.47 (d, 2H, $J=7$ Hz), 3.6 (d, 2H, $J=12$ Hz), 3.75 (d, 3H, $J=11$ Hz), 3.79 (s, 3H), 3.94 (s, 2H), 5.26 (s, 2H), 5.53 (t, 1H, $J=7$ Hz) ppm; ^{31}P (121.4 MHz, CD_3OD) δ 21.5 ppm; MS (m/z) 385.2 $[\text{M}-\text{H}]^+$, 387.1 $[\text{M}+\text{H}]^+$.

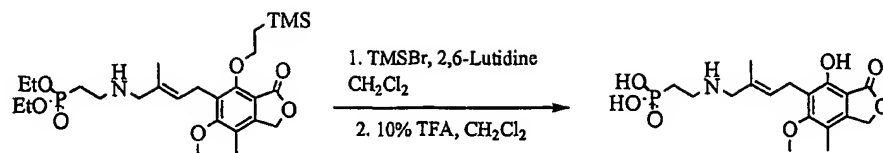
Example 303

Representative compounds of the invention can be prepared as illustrated below.



(2-{4-[6-Methoxy-7-methyl-3-oxo-4-(2-trimethylsilyl-ethoxy)-1,3-dihydro-isobenzofuran-5-yl]-2-methyl-but-2-enylamino}-ethyl)-phosphonic acid diethyl ester

To a solution of 4-[6-methoxy-7-methyl-3-oxo-4-(2-trimethylsilyl-ethoxy)-1,3-dihydro-isobenzofuran-5-yl]-2-methyl-but-2-enal (84 mg, 0.22 mmol), (2-amino-ethyl)-phosphonic acid diethyl ester oxalate (91 mg, 0.33 mmol), and sodium triacetoxyborohydride (93 mg, 0.44 mmol) in DMF (1.5 mL) was added acetic acid (60 μL , 1.0 mmol) at room temperature. The solution was stirred for 2 days when it was quenched by addition of saturated aqueous sodium bicarbonate solution and EtOAc. The organic layer was separated and concentrated under reduced pressure. The residue was purified by RP HPLC using a C18 column with a gradient of H_2O , 0.1% TFA-acetonitrile, 0.1% TFA to provide 115 mg (96%) of the product as an oil. ^1H NMR (300 MHz, CDCl_3) δ 0.04 (s, 9H), 1.16- 1.27 (m, 2H), 1.34 (t, 6H, $J=7$ Hz), 1.94 (s, 3H), 2.18 (s, 3H), 2.20- 2.31 (m, 2H), 3.13- 3.31 (m, 2H), 3.48 (d, 2H, $J=7$ Hz), 3.54 (s, 2H), 3.78 (s, 3H), 4.14 (pent, 4H, $J=7$ Hz), 4.30- 4.37 (m, 2H), 5.13 (s, 2H), 5.65 (t, 1H, $J=7$ Hz), 6.23 (br s, 2H) ppm; ^{31}P (121.4 MHz, CDCl_3) δ 27.8 ppm; MS (m/z) 542.3 $[\text{M}+\text{H}]^+$, 564.2 $[\text{M}+\text{Na}]^+$.

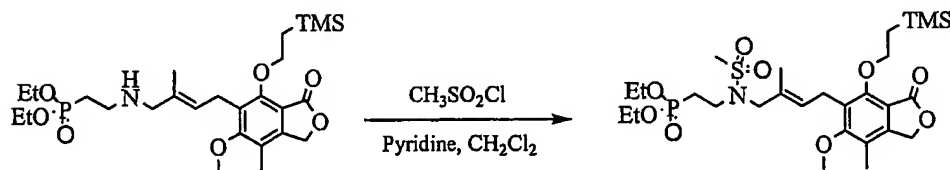


{2-[4-(4-Hydroxy-6-methoxy-7-methyl-3-oxo-1,3-dihydro-isobenzofuran-5-yl)-2-methyl-but-2-enylamino]-ethyl}-phosphonic acid

A solution of (2-{4-[6-methoxy-7-methyl-3-oxo-4-(2-trimethylsilyl-ethoxy)-1,3-dihydro-isobenzofuran-5-yl]-2-methyl-but-2-enylamino}-ethyl)-phosphonic acid diethyl ester (30 mg, 0.055 mmol), TMSBr (72 μ L, 0.55 mmol), and 2,6-lutidine (64 μ L, 0.55 mmol) was stirred in CH_2Cl_2 (1 mL) and DMF (0.5 mL) for 1 hour at ambient temperature. The reaction mixture was purified by RP HPLC using a C18 column with a gradient of H_2O , 0.1% TFA-acetonitrile, 0.1% TFA to provide 7.8 mg of the product as a white solid. ^1H NMR (300 MHz, CD_3OD) δ 1.96 (s, 3H), 1.95- 2.07 (m, 2H), 2.16 (s, 3H), 3.10-3.24 (m, 2H), 3.51 (d, 2H, $J=7$ Hz), 3.57 (s, 2H), 3.81 (s, 3H), 5.25 (s, 2H), 5.73 (t, 1H, $J=7$ Hz) ppm; ^{31}P (121.4 MHz, CD_3OD) δ 20.2 ppm; ^{19}F NMR (282.6 MHz, CD_3OD) δ -74.0 ppm; MS (m/z) 386.3 $[\text{M}+\text{H}]^+$.

Example 304

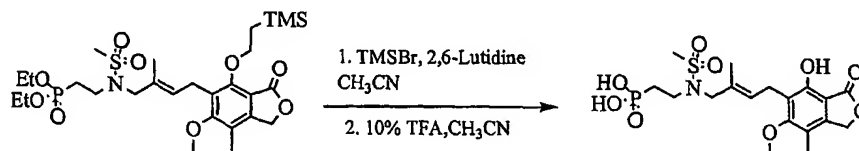
Representative compounds of the invention can be prepared as illustrated below.



[2-(Methanesulfonyl-{4-[6-methoxy-7-methyl-3-oxo-4-(2-trimethylsilyl-ethoxy)-1,3-dihydro-isobenzofuran-5-yl]-2-methyl-but-2-enyl}-amino)-ethyl]-phosphonic acid diethyl ester

A solution of (2-{4-[6-methoxy-7-methyl-3-oxo-4-(2-trimethylsilyl-ethoxy)-1,3-dihydro-isobenzofuran-5-yl]-2-methyl-but-2-enylamino}-ethyl)-

phosphonic acid diethyl ester (45 mg, 0.092 mmol) in CH_2Cl_2 (0.5 mL) was stirred with methanesulfonyl chloride (21 μL , 0.28 mmol) and pyridine (45 μL , 0.55 mmol) at ambient temperature overnight. The reaction was quenched by addition of 2 drops of water. The reaction mixture was concentrated and purified by RP HPLC using a C18 column with a gradient of H_2O , 0.1% TFA-acetonitrile, 0.1% TFA to provide 36 mg of the product (63%) as a clear gel. ^1H NMR (300 MHz, CDCl_3) δ 0.05 (s, 9H), 1.18- 1.29 (m, 2H), 1.29 (t, 6H, $J=7$ Hz), 1.85 (s, 3H), 2.00- 2.13 (m, 2H), 2.19 (s, 3H), 2.85 (s, 3H), 3.32- 3.43 (m, 2H), 3.47 (d, 2H, $J=7$ Hz), 3.69 (s, 2H), 3.79 (s, 3H), 4.05 (pent, 4H, $J=7$ Hz), 4.30- 4.37 (m, 2H), 5.13 (s, 2H), 5.45 (t, 1H, $J=7$ Hz) ppm; ^{31}P (121.4 MHz, CD_3Cl) δ 27.5 ppm; MS (m/z) 642.2 $[\text{M}+\text{Na}]^+$.

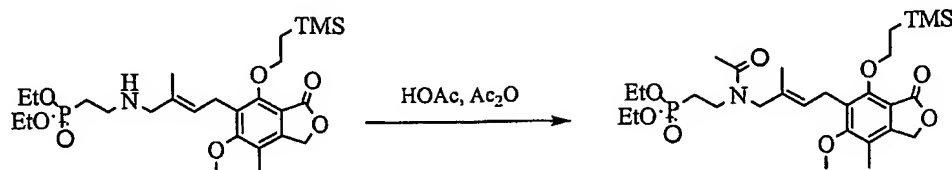


(2-[[4-(4-Hydroxy-6-methoxy-7-methyl-3-oxo-1,3-dihydro-isobenzofuran-5-yl)-2-methyl-but-2-enyl]-methanesulfonyl-amino}-ethyl)-phosphonic acid

A solution of [2-(methanesulfonyl)-{4-[6-methoxy-7-methyl-3-oxo-4-(2-trimethylsilyl-ethoxy)-1,3-dihydro-isobenzofuran-5-yl]-2-methyl-but-2-enyl}-amino]-ethyl]-phosphonic acid diethyl ester (18 mg, 0.029 mmol) in acetonitrile (0.5 mL) was stirred with TMSBr (38 μL , 0.29 mmol) and 2,6-lutidine (34 μL , 0.29 mmol) for 2 hours at room temperature. The reaction was worked up by addition of EtOAc and aqueous 1N HCl. The organic layer was washed with brine and the solvent was removed *in vacuo*. The residue was suspended in a solution of 10% TFA- CH_2Cl_2 for 10 minutes before it was dried to provide 9.9 mg of the desired product (73%) as a white solid. ^1H NMR (300 MHz, $\text{DMSO}-d_6$) δ 1.76 (s, 3H), 1.76- 1.88 (m, 2H), 2.10 (s, 3H), 2.87 (s, 3H), 3.24- 3.35 (m, 2H), 3.39 (d, 2H, $J=7$ Hz), 3.65 (s, 2H), 3.75 (s, 3H), 5.22 (s, 2H), 5.41- 5.48 (m, 1H) ppm; ^{31}P (121.4 MHz, $\text{DMSO}-d_6$) δ 21.4 ppm; MS (m/z) 464.1 $[\text{M}+\text{H}]^+$.

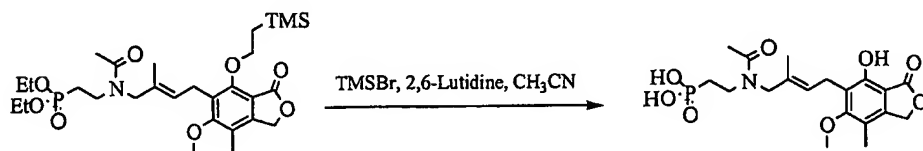
Example 305

Representative compounds of the invention can be prepared as illustrated below.



[2-(Acetyl-{4-[6-methoxy-7-methyl-3-oxo-4-(2-trimethylsilyl-ethoxy)-1,3-dihydro-isobenzofuran-5-yl]-2-methyl-but-2-enyl}-amino)-ethyl]-phosphonic acid diethyl ester

To a solution of (2-{4-[6-methoxy-7-methyl-3-oxo-4-(2-trimethylsilyl-ethoxy)-1,3-dihydro-isobenzofuran-5-yl]-2-methyl-but-2-enylamino}-ethyl)-phosphonic acid diethyl ester (32 mg, 0.059 mmol) in acetic acid (0.5 mL) was added acetic anhydride (0.5 mL). The solution was stirred at room temperature for 90 minutes when it was quenched by addition of 2 drops of water. The solution was dried *in vacuo* and the residue was purified by RP HPLC using a C18 column with a gradient of H₂O, 0.1% TFA-acetonitrile, 0.1% TFA to provide 28 mg of the product (81%) as a clear gel. The NMR data of this compound shows two rotamers in a ratio of 70:30. ¹H NMR (300 MHz, CDCl₃) δ 0.05 (s, 9H), 1.17- 1.27 (m, 2H), 1.30 and 1.31 (t, 6H, *J*= 7 Hz), 1.70-1.79 (m, 2H), 1.76 (s, 3H), 2.00 (s, 3H), 2.18 (s, 3H), 3.40- 3.52 (m, 2H), 3.46 (d, 2H, *J*= 7 Hz), 3.77 (s, 3H), 3.79 and 3.93 (s, 3H), 4.07 (pent, 4H, *J*= 7 Hz), 4.27- 4.35 (m, 2H), 5.13 (s, 2H), 5.22- 5.30 (m, 1H) ppm; ³¹P (121.4 MHz, CDCl₃) δ 27.5 and 28.9 ppm; MS (*m/z*) 584.1 [M+H]⁺, 606.2 [M+Na]⁺.

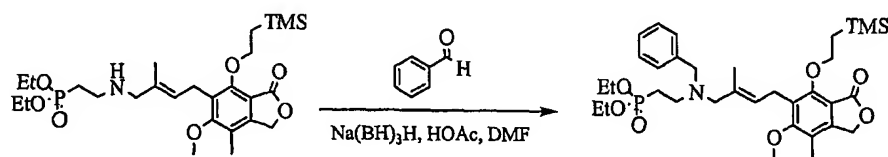


(2-{Acetyl-[4-(4-hydroxy-6-methoxy-7-methyl-3-oxo-1,3-dihydro-isobenzofuran-5-yl)-2-methyl-but-2-enyl]-amino}-ethyl)-phosphonic acid

To a solution of [2-(acetyl-{4-[6-methoxy-7-methyl-3-oxo-4-(2-trimethylsilyl-ethoxy)-1,3-dihydro-isobenzofuran-5-yl]-2-methyl-but-2-enyl}-amino)-ethyl]-phosphonic acid diethyl ester (14 mg, 0.024 mmol) in acetonitrile (0.5 mL) was added TMSBr (31 μ L, 0.24 mmol) and 2,6-lutidine (28 μ L, 0.24 mmol). The solution was stirred at room temperature for 1 hour. The reaction was quenched by addition of methanol and aqueous 1N HCl. The product was extracted with EtOAc. The combined organic extracts were dried over Na₂SO₄ and concentrated *in vacuo*. The product was purified by RP HPLC using a C18 column with a gradient of H₂O, 0.1% TFA-acetonitrile, 0.1% TFA to provide 5.4 mg of the product (53%) as a white solid. The NMR data of this compound shows two rotamers. ¹H NMR (300 MHz, CDCl₃) δ 1.67 and 1.73 (s, 3H), 1.85-2.12 (m, 5H), 2.13 (s, 3H), 3.30-3.61 (m, 4H), 3.75 (s, 3H), 3.76 (br s, 2H), 5.17 (s, 2H), 5.31 (br s, 1H) ppm; ³¹P (121.4 MHz, CDCl₃) δ 27.5 and 28.8 ppm; MS (*m/z*) 428.2 [M+H]⁺, 450.2 [M+Na]⁺.

Example 306

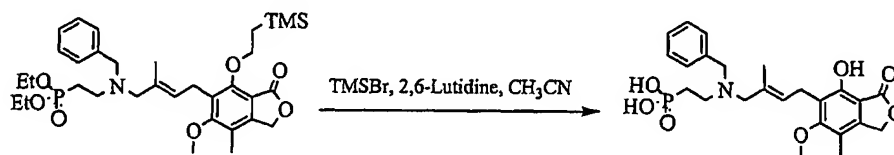
Representative compounds of the invention can be prepared as illustrated below.



[2-(Benzyl-{4-[6-methoxy-7-methyl-3-oxo-4-(2-trimethylsilyl-ethoxy)-1,3-dihydro-isobenzofuran-5-yl]-2-methyl-but-2-enyl}-amino)-ethyl)-phosphonic acid diethyl ester

A solution of (2-{4-[6-methoxy-7-methyl-3-oxo-4-(2-trimethylsilyl-ethoxy)-1,3-dihydro-isobenzofuran-5-yl]-2-methyl-but-2-enylamino}-ethyl)-phosphonic acid diethyl ester (30 mg, 0.055 mmol), benzaldehyde (5.6 μ L, 0.055 mmol), and sodium triacetoxyborohydride (23 mg, 0.11 mmol) was stirred with

acetic acid (15.7 μ L, 0.28 mmol) in DMF (0.5 mL) at room temperature over night. The reaction was quenched with a 10% aqueous Na_2CO_3 solution and the product was extracted with EtOAc. The organic layer was dried and concentrated under reduced pressure. The product was purified by RP HPLC using a C18 column with a gradient of H_2O , 0.1% TFA-acetonitrile, 0.1% TFA to provide 15 mg of the product (43%) as a clear gel. ^1H NMR (300 MHz, CDCl_3) δ 0.02 (s, 9H), 1.18- 1.25 (m, 2H), 1.24 (t, 6H, $J=7$ Hz), 1.86 (s, 3H), 1.88- 2.02 (m, 2H), 2.16 (s, 3H), 2.65- 2.74 (m, 2H), 3.93 (s, 2H), 3.46 (br d, 4H, $J=7$ Hz), 3.76 (s, 3H), 4.00 (pent, 4H, $J=7$ Hz), 4.25- 4.34 (m, 2H), 5.11 (s, 2H), 5.34- 5.43 (m, 1H), 7.18- 7.33 (m, 5H) ppm; ^{31}P (121.4 MHz, CDCl_3) δ 30.9 ppm; MS (m/z) 632.4 $[\text{M}+\text{H}]^+$, 654.3 $[\text{M}+\text{Na}]^+$.

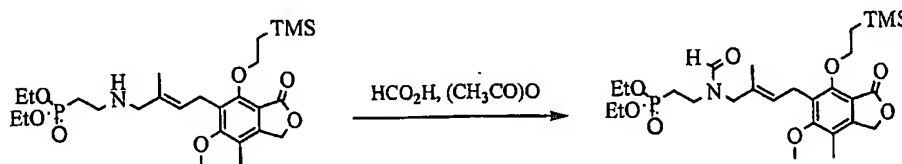


(2-{Benzyl-[4-(4-hydroxy-6-methoxy-7-methyl-3-oxo-1,3-dihydro-isobenzofuran-5-yl)-2-methyl-but-2-enyl]-amino}-ethyl)-phosphonic acid

A solution of (2-{4-[6-methoxy-7-methyl-3-oxo-4-(2-trimethylsilyl-ethoxy)-1,3-dihydro-isobenzofuran-5-yl]-2-methyl-but-2-enylamino}-ethyl)-phosphonic acid diethyl ester (15 mg, 0.024 mmol) in acetonitrile (0.5 mL) was treated with TMSBr (31 μ L, 0.24 mmol) and 2,6-lutidine (28 μ L, 0.24 mmol). The solution was stirred at ambient temperature for 1 hour, when it was quenched with methanol. The solvent was removed under reduced pressure and the residue was purified by RP HPLC using a C18 column with a gradient of H_2O , 0.1% TFA-acetonitrile, 0.1% TFA to provide 11 mg of the product (93%) as a white solid. ^1H NMR (300 MHz, CD_3OD) δ 1.89 (s, 3H), 2.03- 2.15 (m, 2H), 2.14 (s, 3H), 3.30- 3.47 (m, 2H), 3.50 (br s, 2H), 3.62 (br s, 2H), 3.79 (s, 3H), 4.28 (s, 2H), 5.23 (s, 2H), 5.76 (br s, 1H), 7.46 (br s, 5H) ppm; ^{31}P (121.4 MHz, CDCl_3) δ 20.1 ppm; MS (m/z) 476.3 $[\text{M}+\text{H}]^+$, 498.3 $[\text{M}+\text{Na}]^+$.

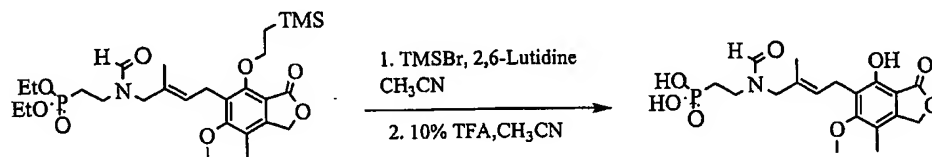
Example 307

Representative compounds of the invention can be prepared as illustrated below.



[2-(Formyl-{4-[6-methoxy-7-methyl-3-oxo-4-(2-trimethylsilanyl-ethoxy)-1,3-dihydro-isobenzofuran-5-yl]-2-methyl-but-2-enyl}-amino)-ethyl]-phosphonic acid diethyl ester

To a solution of (2-{4-[6-Methoxy-7-methyl-3-oxo-4-(2-trimethylsilanyl-ethoxy)-1,3-dihydro-isobenzofuran-5-yl]-2-methyl-but-2-enylamino}-ethyl)-phosphonic acid diethyl ester (74 mg, 0.14 mmol) in formic acid (1 mL) was added formic anhydride (1 mL) and the solution was stirred at room temperature for 1 hour. The reaction mixture was concentrated and the crude product carried onto the next step. The NMR data of this compound shows two rotamers with the ratio of 70:30. ^1H NMR (300 MHz, CDCl_3) δ 0.05 (s, 9H), 1.18- 1.28 (m, 2H), 1.28 and 1.30 (t, 6H, $J=7$ Hz), 1.74 (s, 3H), 1.84- 2.08 (m, 2H), 2.19 (s, 3H), 3.34- 3.45 (m, 2H), 3.47 (d, 2H, $J=7$ Hz), 3.72 and 3.87 (s, 2H), 3.78 and 3.79 (s, 3H), 4.06 and 4.07 (pent, 4H, $J=7$ Hz), 4.26- 4.37 (m, 2H), 5.13 (s, 2H), 5.30-5.46 (m, 1H), 8.03 and 8.19 (s, 1H) ppm; ^{31}P (121.4 MHz, CDCl_3) δ 27.5 and 28.1 ppm; MS (m/z) 570.1 $[\text{M}+\text{H}]^+$, 592.2 $[\text{M}+\text{Na}]^+$.



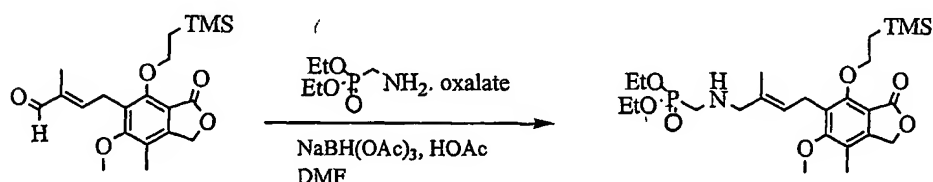
(2-{Formyl-[4-(4-hydroxy-6-methoxy-7-methyl-3-oxo-1,3-dihydro-isobenzofuran-5-yl)-2-methyl-but-2-enyl]-amino}-ethyl)-phosphonic acid

To a solution of crude [2-(formyl-{4-[6-methoxy-7-methyl-3-oxo-4-(2-trimethylsilanyl-ethoxy)-1,3-dihydro-isobenzofuran-5-yl]-2-methyl-but-2-enyl}-

amino)-ethyl]-phosphonic acid diethyl ester (78 mg, 0.14 mmol) in acetonitrile (1 mL) was added TMSBr (177 μ L, 1.4 mmol) and 2,6-lutidine (163 μ L, 1.4 mmol). The solution was stirred at room temperature for 1 hour when it was quenched by addition of methanol and 1N aqueous HCl. The product was extracted with EtOAc and purified by RP HPLC using a C18 column with a gradient of H₂O, 0.1% TFA-acetonitrile, 0.1% TFA to provide 29 mg of the product as a white solid. The NMR data of this compound shows two rotamers with the ratio of approximately 70:30. ¹H NMR (300 MHz, CD₃OD) δ 1.62 and 1.64 (s, 3H), 1.83- 1.98 (m, 2H), 2.16 (s, 3H), 3.38- 3.55 (m, 4H), 3.78 (s, 3H), 3.80 and 3.91 (s, 2H), 5.22 (s, 2H), 5.39- 5.52 (m, 1H), 8.03 and 8.18 (s, 1H) ppm; MS (*m/z*) 414.2 [M+H]⁺, 436.2 [M+Na]⁺.

Example 308

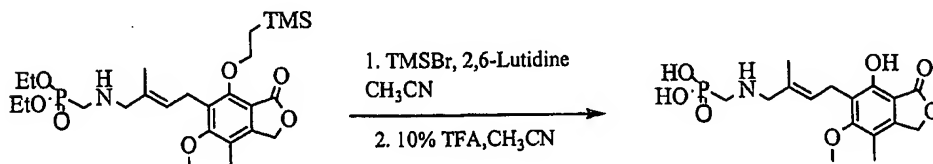
Representative compounds of the invention can be prepared as illustrated below.



({4-[6-Methoxy-7-methyl-3-oxo-4-(2-trimethylsilyl-ethoxy)-1,3-dihydro-isobenzofuran-5-yl]-2-methyl-but-2-enylamino}-methyl)-phosphonic acid diethyl ester

To a solution of 4-[6-methoxy-7-methyl-3-oxo-4-(2-trimethylsilyl-ethoxy)-1,3-dihydro-isobenzofuran-5-yl]-2-methyl-but-2-enal (500 mg, 1.33 mmol), (2-aminomethyl)phosphonic acid diethyl ester oxalate (376 mg, 1.46 mmol), sodium triacetoxyborohydride (563 mg, 2.66 mmol) in DMF (10 mL) was added acetic acid (380 μ L, 6.65 mmol) at room temperature. The solution was stirred overnight when it was quenched by addition of saturated aqueous sodium bicarbonate solution and EtOAc. The organic layer was separated and concentrated under reduced pressure. The residue was purified by silica gel chromatography to provide 500 mg (71%) of the product as an oil. ¹H NMR

(300 MHz, CDCl_3) δ 0.00 (s, 9H), 1.13- 1.23 (m, 2H), 1.25 and 1.27 (t, 6H, $J=7$ Hz), 1.65- 1.75 (m, 2H), 1.77 (s, 3H), 2.13 (s, 3H), 2.80 (s, 1H), 3.14 (s, 2H), 3.41 (d, 2H, $J=7$ Hz), 3.73 (s, 3H), 4.08 and 4.09 (pent, 4H, $J=7$ Hz), 4.20- 4.30 (m, 2H), 5.08 (s, 2H), 5.30 (t, 1H, $J=7$ Hz) ppm; ^{31}P (121.4 MHz, CDCl_3) δ 26.5 ppm; MS (m/z) 528.1 $[\text{M}+\text{H}]^+$, 550.2 $[\text{M}+\text{Na}]^+$.

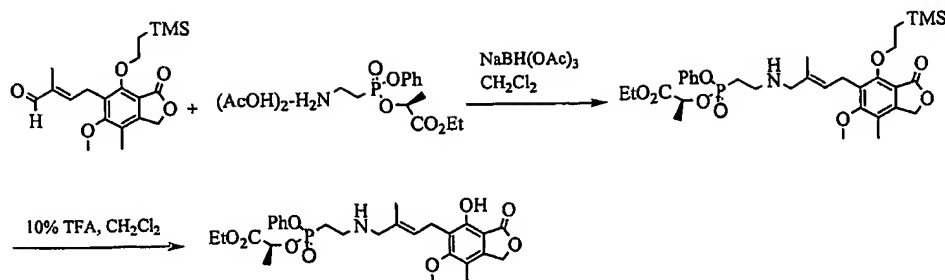


{[4-(4-Hydroxy-6-methoxy-7-methyl-3-oxo-1,3-dihydro-isobenzofuran-5-yl)-2-methyl-but-2-enylamino]-methyl}-phosphonic acid

To a solution of ({4-[6-methoxy-7-methyl-3-oxo-4-(2-trimethylsilyl-ethoxy)-1,3-dihydro-isobenzofuran-5-yl]-2-methyl-but-2-enylamino}-methyl)-phosphonic acid diethyl ester (20 mg, 0.038 mmol) in DMF (0.5 mL) was added TMSBr (49 μL , 0.38 mmol) and 2,6-lutidine (44 μL , 0.38 mmol). The solution was stirred at room temperature for 1 hour when it was quenched by addition of methanol. The product was purified by RP HPLC using a C18 column with a gradient of H_2O , 0.1% TFA-acetonitrile, 0.1% TFA to provide 5.6 mg of the product as a white solid. ^1H NMR (300 MHz, CD_3OD and CDCl_3) δ 1.93 (s, 3H), 2.13 (s, 3H), 2.94 (br d, 2H, $J=11$ Hz), 3.42-3.53 (m, 2H), 3.60 (s, 2H), 3.78 (s, 3H), 5.22 (s, 2H), 5.71 (br s, 1H) ppm; ^{31}P (121.4 MHz, CDCl_3) δ 8.5 ppm; MS (m/z) 372.2 $[\text{M}+\text{H}]^+$, 743.2 $[2\text{M}+\text{H}]^+$.

Example 309

Representative compounds of the invention can be prepared as illustrated below.

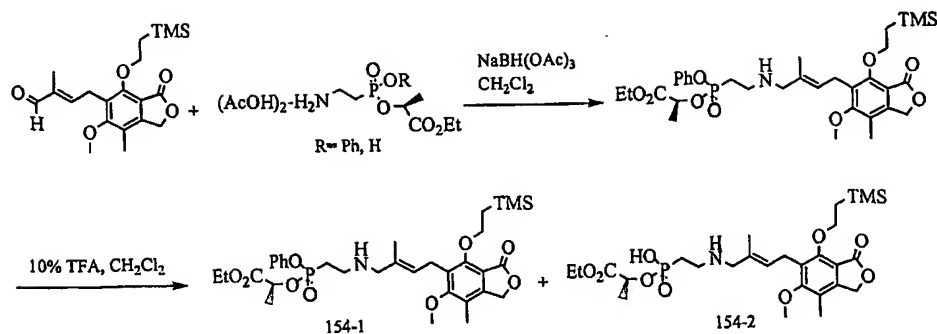


2-([4-(4-Hydroxy-6-methoxy-7-methyl-3-oxo-1,3-dihydro-isobenzofuran-5-yl)]-2-methyl-but-2-enylamino)-ethyl-phenoxy-phosphinoyloxy)-propionic acid ethyl ester

A solution of 4-[6-methoxy-7-methyl-3-oxo-4-(2-trimethylsilanyloxy)-1,3-dihydro-isobenzofuran-5-yl]-2-methyl-but-2-enal (188 mg, 0.5 mmol) was stirred with 2-[(2-aminoethyl)phenoxy-phosphinoyloxy]-propionic acid ethyl ester acetic acid salt (315.8 mg, 0.75 mmol) in CH_2Cl_2 (3 mL) for 2 hours at ambient temperature. Sodium triacetoxyborohydride (159 mg, 0.75 mmol) was added to the solution and the reaction was allowed to proceed for 1 hour. The reaction was quenched by addition of a saturated aqueous solution of NaHCO_3 and the product was extracted with EtOAc. The organic layer was removed under reduced pressure and the residue was resuspended in a 10% TFA/ CH_2Cl_2 for 1 hour. The reaction mixture was concentrated and the product was purified by RP HPLC using a C18 column with a gradient of H_2O , 0.1% TFA-acetonitrile, 0.1% TFA to provide 198 mg of the product as a white solid. The NMR data of this compound shows two diastereomers at phosphorus in a ratio of approximately 45: 55. ^1H NMR (300 MHz, CD_3OD) δ 1.23 and 1.24 (t, 3H, $J=7$ Hz), 1.38 and 1.52 (d, 3H, $J=7$ Hz), 1.97 and 1.98 (s, 3H), 2.14 (s, 3H), 2.44- 2.66 (m, 2H), 3.31- 3.48 (m, 2H), 3.51 (d, 2H, $J=7$ Hz), 3.66 (d, 2H, $J=5$ Hz), 3.80 (s, 3H), 4.10- 4.27 (m, 2H), 4.90- 5.10 (m, 1H), 5.20 (s, 2H), 5.73- 5.82 (m, 1H), 7.15- 7.27 (m, 3H), 7.35- 7.45 (m, 2H) ppm; ^{31}P (121.4 MHz, CD_3OD) δ 22.6, 24.3 ppm; MS (m/z) 561.9 $[\text{M}+\text{H}]^+$.

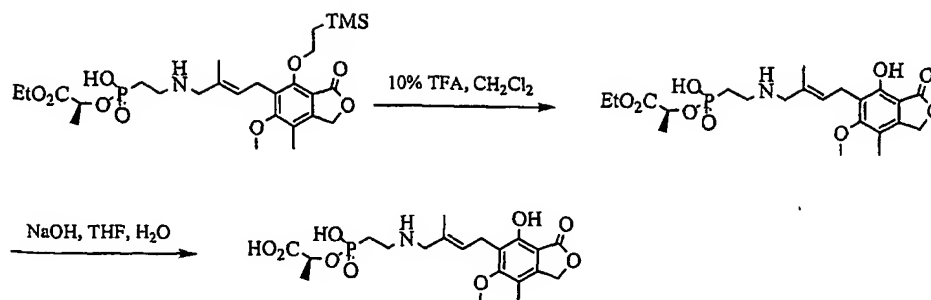
Example 310

Representative compounds of the invention can be prepared as illustrated below.



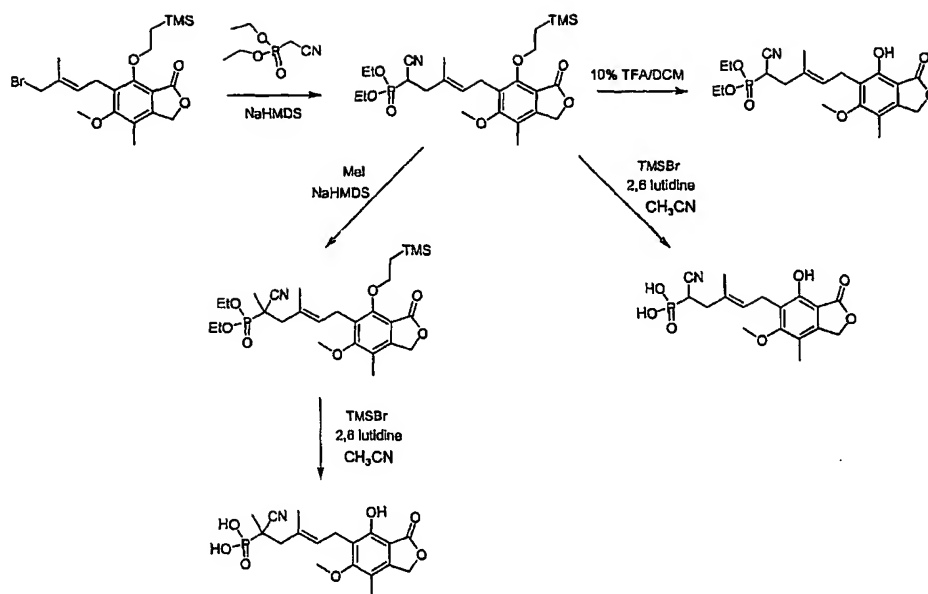
2-[Hydroxy-(2-{4-[6-methoxy-7-methyl-3-oxo-4-(2-trimethylsilyl-ethoxy)-1,3-dihydro-isobenzofuran-5-yl]-2-methyl-but-2-enylamino}-ethyl)-phosphinoyloxy]-propionic acid ethyl ester

A solution of 4-[6-methoxy-7-methyl-3-oxo-4-(2-trimethylsilyl-ethoxy)-1,3-dihydro-isobenzofuran-5-yl]-2-methyl-but-2-enal (38 mg, 0.1 mmol) was stirred with 2-[(2-aminoethyl)-phenoxy-phosphinoyloxy]-propionic acid ethyl ester acetic acid (63 mg, 0.15 mmol) in CH_2Cl_2 (1 mL) for 2 hours at ambient temperature. Sodium triacetoxyborohydride (32 mg, 0.15 mmol) was added to the solution and the reaction was allowed to proceed for 1 hour. The reaction was quenched by addition of a saturated aqueous solution of NaHCO_3 and the product was extracted with EtOAc. The organic layer was removed under reduced pressure and the residue was re-suspended in 10% TFA/ CH_2Cl_2 for 1 hour. The reaction mixture was concentrated and the product was purified by RP HPLC using a C18 column with a gradient of H_2O , 0.1% TFA-acetonitrile, 0.1% TFA to provide 15 mg of the product (154-2). ^1H NMR (300 MHz, CDCl_3) δ 0.04 (s, 9H), 1.15- 1.24 (m, 2H), 1.26 (t, 3H, $J = 7$ Hz), 1.48 (d, 3H, $J = 7$ Hz), 1.93 (s, 3H), 2.10- 2.25 (m, 2H), 2.18 (s, 3H), 3.10- 3.31 (m, 2H), 3.48 (d, 2H, $J = 7$ Hz), 3.48- 3.61 (m, 2H), 3.77 (s, 3H), 4.04- 4.21 (m, 2H), 4.29- 4.40 (m, 2H), 4.81- 4.92 (m, 1H), 5.13 (s, 2H), 5.64 (t, 1H, $J = 7$ Hz), 8.70- 9.11 (m, 3H) ppm; ^{31}P (121.4 MHz, CDCl_3) δ 21.9 ppm; MS (m/z) 586.3 $[\text{M}+\text{H}]^+$, 1171.4 $[2\text{M}+\text{H}]^+$.

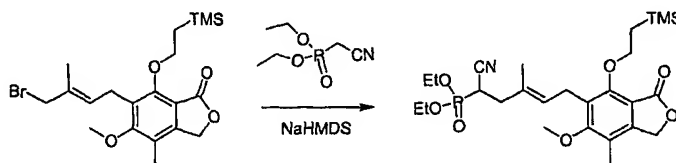


2-(Hydroxy-{2-[4-(4-hydroxy-6-methoxy-7-methyl-3-oxo-1,3-dihydro-isobenzofuran-5-yl)-2-methyl-but-2-enylamino]-ethyl}-phosphinoyloxy)-propionic acid

A solution of 2-[hydroxy-(2-{4-[6-methoxy-7-methyl-3-oxo-4-(2-trimethylsilyl-ethoxy)-1,3-dihydro-isobenzofuran-5-yl]-2-methyl-but-2-enylamino}-ethyl)-phosphinoyloxy]-propionic acid ethyl ester (15 mg, 0.026 mmol) in 10% TFA-CH₂Cl₂ (1 mL) was stirred at ambient temperature for 10 minutes. The reaction was worked up by removal of the solvent. The residue was dissolved in THF (0.5 mL) and water (0.4 mL) and 1N aqueous NaOH solution (0.1 mL) was added. The solution was stirred at room temperature for 20 minutes when it was acidified with 1N aqueous HCl solution. The resulting solution was purified by RP HPLC using a C18 column with a gradient of H₂O, 0.1% TFA-acetonitrile, 0.1% TFA to provide 6.8 mg of the product as a white solid. ¹H NMR (300 MHz, CDCl₃) δ 1.38 (d, 3H, *J*= 7 Hz), 1.91 (s, 3H), 2.13 (s, 3H), 2.12- 2.28 (m, 2H), 3.12- 3.33 (m, 2H), 3.41 (d, 2H, *J*= 6 Hz), 3.56 (br s, 2H), 3.75 (s, 3H), 4.71- 4.88 (m, 1H), 5.16 (s, 2H), 5.58- 5.71 (m, 1H), 7.88 (br s, 3H), 8.60 (br s, 1H), 8.78 (br s, 1H) ppm; ³¹P (121.4 MHz, CDCl₃) δ 22.0 ppm; MS (*m/z*) 458.3 [M+H]⁺, 480.3 [M+Na]⁺.

**Example 311**

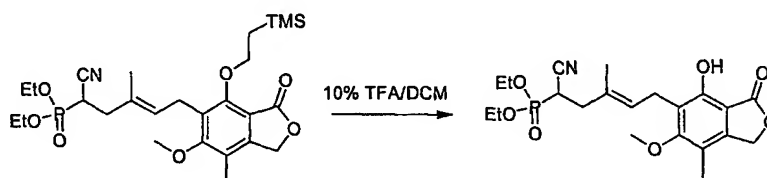
Representative compounds of the invention can be prepared as illustrated below.



{1-Cyano-5-[6-methoxy-7-methyl-3-oxo-4-(2-trimethylsilylanyl-ethoxy)-1,3-dihydro-isobenzofuran-5-yl]-3-methyl-pent-3-enyl}-phosphonic acid diethyl ester

To a solution of diethyl cyanomethylphosphonate (241 mg, 1.38 mmol) in THF (1 mL) was added a THF solution of sodium bis(trimethylsilyl)amide (1.0 M, 1.13 mL, 1.15 mmol). After stirring for 30 minutes, the solution was added dropwise to a solution of 6-(4-bromo-3-methylbut-2-enyl)-7-hydroxy-5-methoxy-4-methyl-3H-isobenzofuran-1-one (100 mg, 0.23 mmol) in THF (1 mL). The resulting mixture was allowed to stir at room temperature for one hour before saturated aqueous ammonium chloride was added. The reaction mixture was extracted with ethyl acetate. The organic layer was dried over sodium

sulfate and concentrated to dryness. The residue was purified by silica gel column chromatography, affording 110 mg (90 %) of the desired product. ^1H NMR (300 MHz, CDCl_3) δ 0.04 (s, 9H), 1.24 (dd, $J = 7, 8$ Hz, 2H), 1.36 (t, 6H), 1.86 (s, 3H), 2.17 (s, 3H), 2.43-2.57 (m, 2H), 3.04-3.17 (m, 1H), 3.47 (d, $J = 7.2$ Hz, 2H), 3.79 (s, 3H), 4.12-4.37 (m, 6H), 5.13 (s, 2H), 5.44 (t, $J = 7.2$ Hz, 1H) ppm; ^{31}P (121.4 MHz, CDCl_3) δ 18.18 ppm; MS (m/z) 560 $[\text{M}+\text{Na}]^+$.

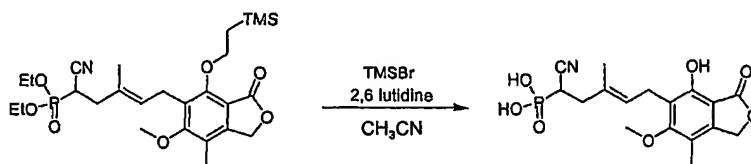


[1-Cyano-5-(4-hydroxy-6-methoxy-7-methyl-3-oxo-1,3-dihydro-isobenzofuran-5-yl)-3-methyl-pent-3-enyl]-phosphonic acid diethyl ester

{1-Cyano-5-[6-methoxy-7-methyl-3-oxo-4-(2-trimethylsilyl-ethoxy)-1,3-dihydro-isobenzofuran-5-yl]-3-methyl-pent-3-enyl}-phosphonic acid diethyl ester (25mg, 0.047 mmol) was dissolved in a solution of 10% TFA/ CH_2Cl_2 (5 mL) and stirred at room temperature for 2 hours. The reaction mixture was dried under reduced pressure and the product was purified by RP-HPLC to provide 16 mg (80%) of the desired product as a white solid. ^1H NMR (300 MHz, CDCl_3) δ 1.38 (t, 6H), 1.86 (s, 3H), 2.15 (s, 3H), 2.40-2.58 (m, 2H), 3.01-3.14 (m, 1H), 3.45 (d, $J = 7.2$ Hz, 2H), 3.79 (s, 3H), 4.18-4.30 (m, 4H), 5.21 (s, 2H), 5.48 (t, $J = 7.2$ Hz, 1H) ppm; ^{31}P (121.4 MHz, CDCl_3) δ 18.09 ppm; MS (m/z) 436 $[\text{M}-\text{H}]^-$, 438 $[\text{M}+\text{H}]^+$.

Example 312

Representative compounds of the invention can be prepared as illustrated below.

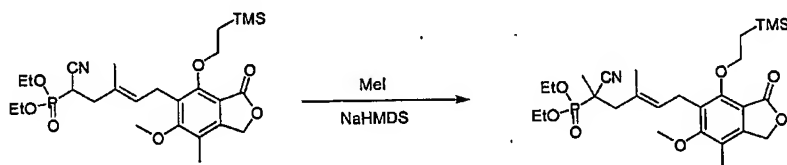


[1-Cyano-5-(4-hydroxy-6-methoxy-7-methyl-3-oxo-1,3-dihydro-isobenzofuran-5-yl)-3-methyl-pent-3-enyl]-phosphonic acid

To a solution of {1-cyano-5-[6-methoxy-7-methyl-3-oxo-4-(2-trimethylsilyl-ethoxy)-1,3-dihydro-isobenzofuran-5-yl]-3-methyl-pent-3-enyl}-phosphonic acid diethyl ester (35 mg, 0.065 mmol) in acetonitrile (2 mL) was added TMSBr (180 μ L, 1.38 mmol) and 2,6-lutidine (160 μ L, 1.38 mmol). The reaction solution was allowed stir at room temperature for one hour before quenching with MeOH. The reaction mixture was dried under reduced pressure and the residue was purified by RP HPLC using a C18 column with a gradient of H₂O, 0.1% TFA-acetonitrile, 0.1% TFA to provide 15 mg (60 %) of the desired product. ¹H NMR (300 MHz, CD₃OD) δ 1.86 (s, 3H), 2.15 (s, 3H), 2.38-2.57 (m, 2H), 3.17-3.28 (m, 1H), 3.44 (d, J = 7.2 Hz, 2H), 3.80 (s, 3H), 5.25 (s, 2H), 5.47 (t, J = 7.2 Hz, 1H) ppm; ³¹P (121.4 MHz, CD₃OD) δ 15.28 ppm; MS (m/z) 380 [M-H]⁻, 382 [M+H]⁺.

Example 313

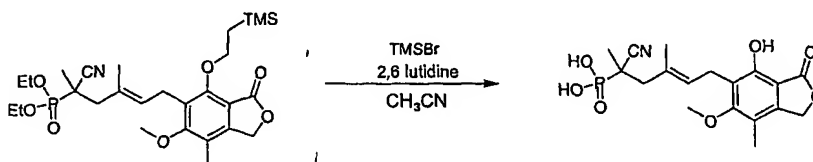
Representative compounds of the invention can be prepared as illustrated below.



{1-Cyano-5-[6-methoxy-7-methyl-3-oxo-4-(2-trimethylsilyl-ethoxy)-1,3-dihydro-isobenzofuran-5-yl]-1,3-dimethyl-pent-3-enyl}-phosphonic acid diethyl ester

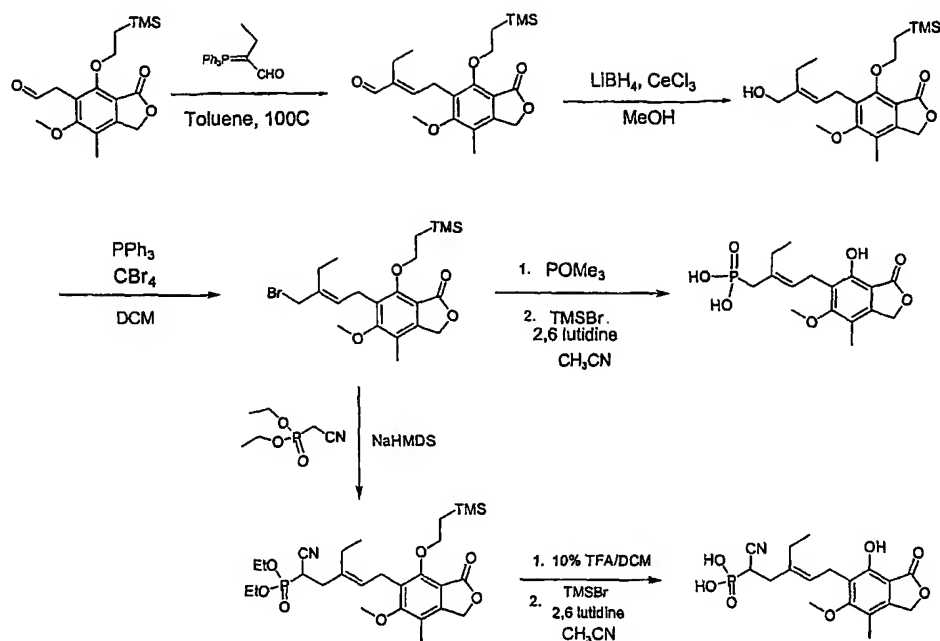
To a solution of {1-cyano-5-[6-methoxy-7-methyl-3-oxo-4-(2-trimethylsilyl-ethoxy)-1,3-dihydro-isobenzofuran-5-yl]-3-methyl-pent-3-enyl}-phosphonic acid diethyl ester (45 mg, 0.084 mmol) in THF (0.5 mL) was added sodium bis(trimethylsilyl)amide (1.0 M, 1.13 mL, 1.15 mmol). After stirring for 20 minutes, iodomethane (52 μ L, 0.84 mmol) was added dropwise

and the resulting mixture was allowed to stir at room temperature for 2 hours. The reaction mixture was quenched with saturated aqueous ammonium chloride and extracted with ethyl acetate. The organic layer was dried over sodium sulfate and concentrated to dryness. The residue was purified by RP HPLC using a C18 column with a gradient of H₂O, 0.1% TFA-acetonitrile, 0.1% TFA to afford 6.6 mg (23 %) of the desired product. ¹H NMR (300 MHz, CDCl₃) δ 0.00 (s, 9H), 1.16 (dd, *J* = 7, 8 Hz, 2H), 1.31 (t, 6H), 1.38 (d, 3H), 1.92 (s, 3H), 2.17 (s, 3H), 2.23 (m, 1H), 2.65 (m, 1H), 3.30-3.42 (m, 2H), 3.73 (s, 3H), 4.14-4.27 (m, 6H), 5.08 (s, 2H), 5.28 (t, *J* = 7.2 Hz, 1H) ppm; ³¹P (121.4 MHz, CDCl₃) δ 22.26 ppm; MS (*m/z*) 574 [M+Na]⁺.



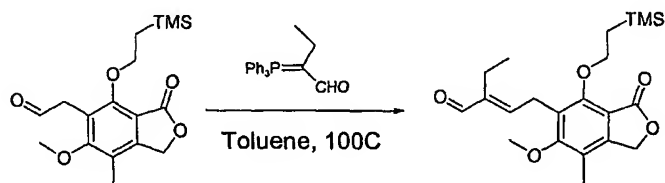
[1-Cyano-5-(4-hydroxy-6-methoxy-7-methyl-3-oxo-1,3-dihydro-isobenzofuran-5-yl)-1,3-dimethyl-pent-3-enyl]-phosphonic acid

To a solution of {1-cyano-5-[6-methoxy-7-methyl-3-oxo-4-(2-trimethylsilylanyl-ethoxy)-1,3-dihydro-isobenzofuran-5-yl]-1,3-dimethyl-pent-3-enyl}-phosphonic acid diethyl ester (18 mg, 0.04 mmol) in DMF (0.5 mL) and DCM (0.5 mL) was added TMSBr (51 μL, 0.4 mmol) and 2,6-lutidine (46 μL, 0.4 mmol). The reaction solution was allowed stir at room temperature overnight before quenching with MeOH. The reaction mixture was dried under reduced pressure and the residue was purified by RP HPLC using a C18 column with a gradient of H₂O, 0.1% TFA-acetonitrile, 0.1% TFA to provide 4.5 mg (33 %) of the desired product. ¹H NMR (300 MHz, CD₃OD) δ 1.37 (d, 3H), 1.87 (s, 3H), 2.13 (s, 3H), 2.26 (m, 1H), 2.64 (m, 1H), 3.39 (m, 2H), 3.75 (s, 3H), 5.18 (s, 2H), 5.34 (m, 1H) ppm; ³¹P (121.4 MHz, CD₃OD) δ 21.47 ppm; MS (*m/z*) 422 [M-H]⁻, 424 [M+H]⁺.



Example 314

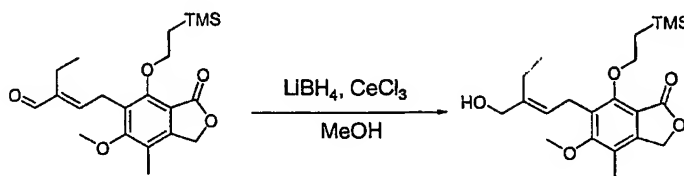
Representative compounds of the invention can be prepared as illustrated below.



2-Ethyl-4-[6-methoxy-7-methyl-3-oxo-4-(2-trimethylsilyl-ethoxy)-1,3-dihydro-isobenzofuran-5-yl]-but-2-enal

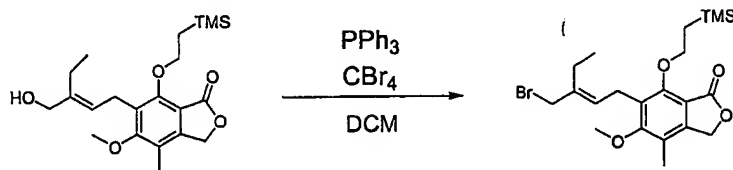
A solution of [6-methoxy-7-methyl-3-oxo-4-(2-trimethylsilyl-ethoxy)-1,3-dihydro-isobenzofuran-5-yl]-acetaldehyde (1.5 g, 4.46 mmol) in toluene (14 mL) was heated at 100 °C with 2-(triphenyl-phosphanylidene)-butyraldehyde (1.68g, 5.35 mmol) overnight. A second portion of 2-(triphenyl-phosphanylidene)-butyraldehyde (495 mg, 1.49 mmol) was added and the reaction mixture was heated for an additional day. After concentration, the residue was purified by silica gel chromatography to provide 1.3g (83%) of the desired product as oil. ¹H NMR (300 MHz, CDCl₃) δ 0.01 (s, 9H), 1.03 (t, 3H),

1.10- 1.21 (m, 2H), 2.15 (s, 3H), 2.15-2.44 (m, 2H), 3.67- 3.76 (m, 2H), 3.74 (s, 3H), 4.31- 4.36 (m, 2H), 5.10 (s, 2H), 6.34- 6.38 (m, 1H), 9.28 (s, 1H) ppm.



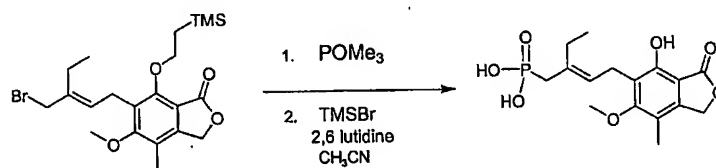
6-(3-Hydroxymethyl-pent-2-enyl)-5-methoxy-4-methyl-7-(2-trimethylsilyl-ethoxy)-3H-isobenzofuran-1-one

A solution of 2-ethyl-4-[6-methoxy-7-methyl-3-oxo-4-(2-trimethylsilyl-ethoxy)-1,3-dihydro-isobenzofuran-5-yl]-but-2-enal (1.3g, 3.30 mmol) in methanol (10 mL) and THF (10 mL) was cooled to 0 °C. A solution of CeCl_3 (8.25 mL, 0.4M, MeOH: H_2O , 9:1) was added, followed by LiBH_4 (1.66 mL, 3.30 mmol of a 2M solution in THF). The ice bath was removed and the reaction mixture was allowed to warm to room temperature. The reaction mixture was stirred for an additional 40 minutes, whereupon TLC indicated complete consumption of starting aldehyde. The reaction was worked up by addition of aqueous 1N HCl and the product was extracted with EtOAc. The organic layer was washed with saturated aqueous sodium bicarbonate solution and brine. The organic layer was concentrated under reduced pressure and the residue was purified by silica gel chromatography to provide 948 mg (73%) of the product as colorless oil. ^1H NMR (300 MHz, CDCl_3) δ 0.00 (s, 9H), 1.07 (t, 3H), 1.20 (dd, 2H, $J = 7, 8$ Hz), 2.13 (s, 3H), 2.38- 2.50 (m, 2H), 3.77 (s, 3H), 3.99 (s, 2H), 4.27 (dd, 2H, $J = 7, 8$ Hz), 5.08 (s, 2H), 5.34 (t, $J = 7.2$ Hz, 1H) ppm.



6-(3-Bromomethyl-pent-2-enyl)-5-methoxy-4-methyl-7-(2-trimethylsilyl-ethoxy)-3H-isobenzofuran-1-one

Polymer-supported triphenylphosphine (3 mmol/g, 0.66 g) was soaked in dichloromethane (6 mL) for 1 hour 6-(3-Hydroxymethyl-pent-2-enyl)-5-methoxy-4-methyl-7-(2-trimethylsilanyl-ethoxy)-3H-isobenzofuran-1-one (260 mg, 0.66 mmol) and carbon tetrabromide (657 mg, 1.98 mmol) were added sequentially and the mixture was shaken for 1 hour at room temperature. The mixture was filtered and the filtrate was concentrated. The residue was purified by silica gel chromatography to provide 233 mg (77%) of the product as a white solid. ^1H NMR (300 MHz, CDCl_3) δ 0.00 (s, 9H), 1.08 (t, 3H), 1.20 (dd, 2H, $J=7, 8$ Hz), 2.14 (s, 3H), 2.35-2.43 (m, 2H), 3.44 (d, $J=7.2$, 2H), 3.73 (s, 3H), 3.95 (s, 2H), 4.27 (dd, 2H, $J=7, 8$ Hz), 5.08 (s, 2H), 5.53 (t, $J=7.2$ Hz, 1H) ppm.

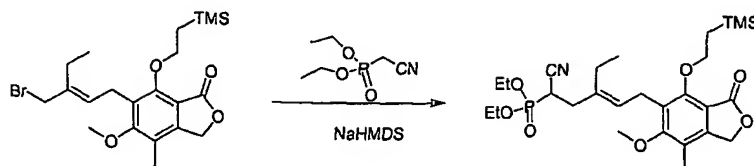


[2-Ethyl-4-(4-hydroxy-6-methoxy-7-methyl-3-oxo-1,3-dihydro-isobenzofuran-5-yl)-but-2-enyl]-phosphonic acid

A solution of 6-(3-bromomethyl-pent-2-enyl)-5-methoxy-4-methyl-7-(2-trimethylsilanyl-ethoxy)-3H-isobenzofuran-1-one (230 mg, 0.5 mmol) in trimethylphosphite (1.5 mL, 12.75 mmol) was heated to 100 °C for 4 hours. The reaction was worked up by removal of excess trimethylphosphite under reduced pressure. The residue was dissolved in acetonitrile (1 mL) and TMSBr (646 μL , 5.0 mmol) and 2,6-lutidine (580 μL , 5.0 mmol) were added at 0 °C. The reaction solution was allowed to warm to room temperature and stirred for 4 hours. The reaction was cooled to 0 °C and quenched with addition of MeOH. The reaction mixture was dried under reduced pressure and the residue was purified by RP HPLC using a C18 column with a gradient of H_2O , 0.1% TFA-acetonitrile, 0.1% TFA to provide 77mg (58%) of the product. ^1H NMR (300 MHz, CD_3OD) δ 1.08 (t, 3H), 2.16 (s, 3H), 2.43 (m, 2H), 2.48 (d, 2H, $J=22$ Hz), 3.46 (t, 2H, $J=6$ Hz), 3.79 (s, 3H), 5.25 (s, 2H), 5.38 (q, 1H, $J=7$ Hz) ppm.; ^{31}P (121.4 MHz, CD_3OD) δ 25.65 ppm.; MS (m/z) 355 [$\text{M}-\text{H}$] $^-$, 357 [$\text{M}+\text{H}$] $^+$.

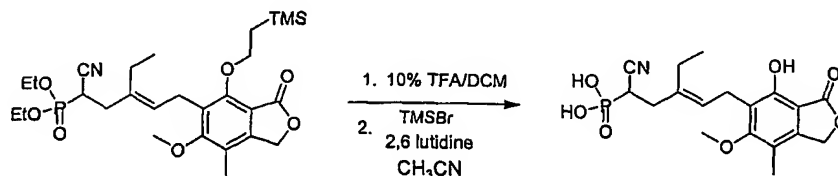
Example 315

Representative compounds of the invention can be prepared as illustrated below.



{1-Cyano-3-ethyl-5-[6-methoxy-7-methyl-3-oxo-4-(2-trimethylsilyl-ethoxy)-1,3-dihydro-isobenzofuran-5-yl]-pent-3-enyl}-phosphonic acid diethyl ester

To a solution of diethyl cyanomethylphosphonate (233 mg, 1.32 mmol) in THF (1 mL) was added a THF solution of sodium bis(trimethylsilyl)amide (1.0 M, 1.21 mL, 1.21 mmol). After stirring for 30 minutes, the solution was added dropwise to a solution of 6-(3-bromomethyl-pent-2-enyl)-5-methoxy-4-methyl-7-(2-trimethylsilyl-ethoxy)-3H-isobenzofuran-1-one (100 mg, 0.22 mmol) in THF (1 mL). The resulting mixture was allowed to stir at room temperature overnight before saturated aqueous ammonium chloride was added. The reaction mixture was extracted with ethyl acetate. The organic layer was dried over sodium sulfate and concentrated to dryness. The residue was purified by preparative reverse-phase HPLC, affording 51 mg (42%) of the desired product. ^1H NMR (300 MHz, CDCl_3) δ 0.04 (s, 9H), 1.07 (t, 3H), 1.24 (dd, 2H, $J = 7, 8$ Hz), 1.36 (t, 6H), 2.12 (m, 1H), 2.18 (s, 3H), 2.35-2.47 (m, 2H), 2.67 (m, 1H), 3.00-3.14 (m, 1H), 3.44 (d, $J = 7.2$, 2H), 3.79 (s, 3H), 4.12-4.37 (m, 6H), 5.13 (s, 2H), 5.38 (t, $J = 7.2$ Hz, 1H) ppm; ^{31}P (121.4 MHz, CDCl_3) δ 18.26 ppm; MS (m/z) 574 $[\text{M}+\text{Na}]^+$.

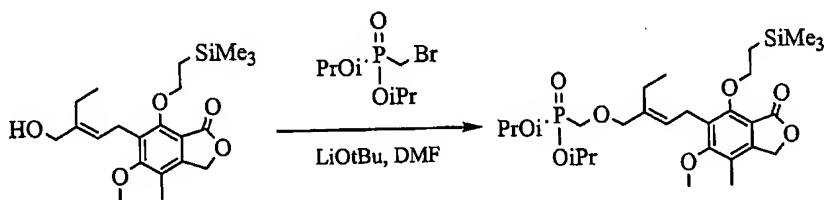


[1-Cyano-3-ethyl-5-(4-hydroxy-6-methoxy-7-methyl-3-oxo-1,3-dihydro-isobenzofuran-5-yl)-pent-3-enyl]-phosphonic acid

{1-Cyano-3-ethyl-5-[6-methoxy-7-methyl-3-oxo-4-(2-trimethylsilyl-ethoxy)-1,3-dihydro-isobenzofuran-5-yl]-pent-3-enyl}-phosphonic acid diethyl ester (19.5 mg, 0.035 mmol) was dissolved in a solution of 10% TFA/ CH₂Cl₂ (2 mL) and stirred at room temperature for 10 minutes. The reaction mixture was dried under reduced pressure and purified by RP-HPLC to provide 9.5 mg (61%) of the desired product. This material was dissolved in DMF (0.5 mL) and DCM (0.5 mL) and TMSBr (27 μ L, 0.2 mmol) and 2,6-lutidine (23 μ L, 0.2 mmol) were added. The reaction solution was allowed stir at room temperature overnight before quenching with MeOH. The reaction mixture was dried under reduced pressure and the residue was purified by RP HPLC using a C18 column with a gradient of H₂O, 0.1% TFA-acetonitrile, 0.1% TFA to provide 5.1 mg (65%) of the desired product as a white solid. ¹H NMR (300 MHz, CD₃OD) δ 1.10 (t, 3H), 2.16 (s, 3H), 2.23-2.52 (m, 3H), 2.67 (m, 1H), 3.05-3.20 (m, 1H), 3.48 (d, J = 7.2, 2H), 3.81 (s, 3H), 5.26 (s, 2H), 5.43 (t, J = 7.2 Hz, 1H) ppm; ³¹P (121.4 MHz, CD₃OD) δ 14.18 ppm; MS (m/z) 394 [M-H]⁻, 396 [M+H]⁺.

Example 316

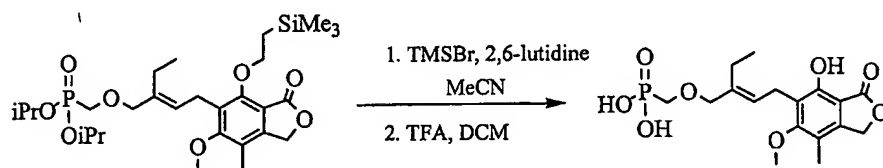
Representative compounds of the invention can be prepared as illustrated below.



{2-Ethyl-4-[6-methoxy-7-methyl-3-oxo-4-(2-trimethylsilyl-ethoxy)-1,3-dihydro-isobenzofuran-5-yl]-but-2-enyloxymethyl}-phosphonic acid diisopropyl ester

To a solution of bromomethylphosphonate diisopropyl ester (680mg, 2.62 mmol) and 6-(3-hydroxymethyl-pent-2-enyl)-5-methoxy-4-methyl-7-(2-trimethylsilyl-ethoxy)-3H-isobenzofuran-1-one (688mg, 1.75 mmol) in DMF

(3 mL) was added lithium t-butoxide (1.0M in THF; 2.6mL). The reaction was heated at 70°C for 2 hours. After cooling to ambient temperature, more bromomethylphosphonate diisopropyl ester (680mg, 2.62 mmol) and lithium t-butoxide (1.0M in THF; 2.6mL) were added. The reaction mixture was heated at 70°C for a further hour, cooled, poured into a solution of lithium chloride (5% aqueous) and extracted with ethyl acetate. The organic extract was dried and the product was purified by chromatography on silica gel, eluting with hexane-ethyl acetate to provide 347 mg (35%) of the product as a colorless oil. ¹H NMR (300 MHz, CDCl₃) δ 0.04 (s, 9H), 1.09 (t, 3H, *J*= 7.5 Hz), 1.20- 1.26 (m, 2H), 1.31 (t, 12H, *J*= 6 Hz), 2.18 (s, 3H), 2.29 (q, 2H, *J*= 7.5 Hz), 3.5 (m, 2H), 3.59 (d, 2H, *J*= 8.7 Hz), 3.78 (s, 3H), 3.98 (s, 2H), 4.28- 4.35 (m, 2H), 4.6-4.8 (m, 2H), 5.13 (s, 2H), 5.4 (t, 1H, *J*= 7 Hz) ppm; ³¹P (121.4 MHz, CDCl₃) δ 20.26 ppm; MS (*m/z*) 593.3 [M+Na]⁺.



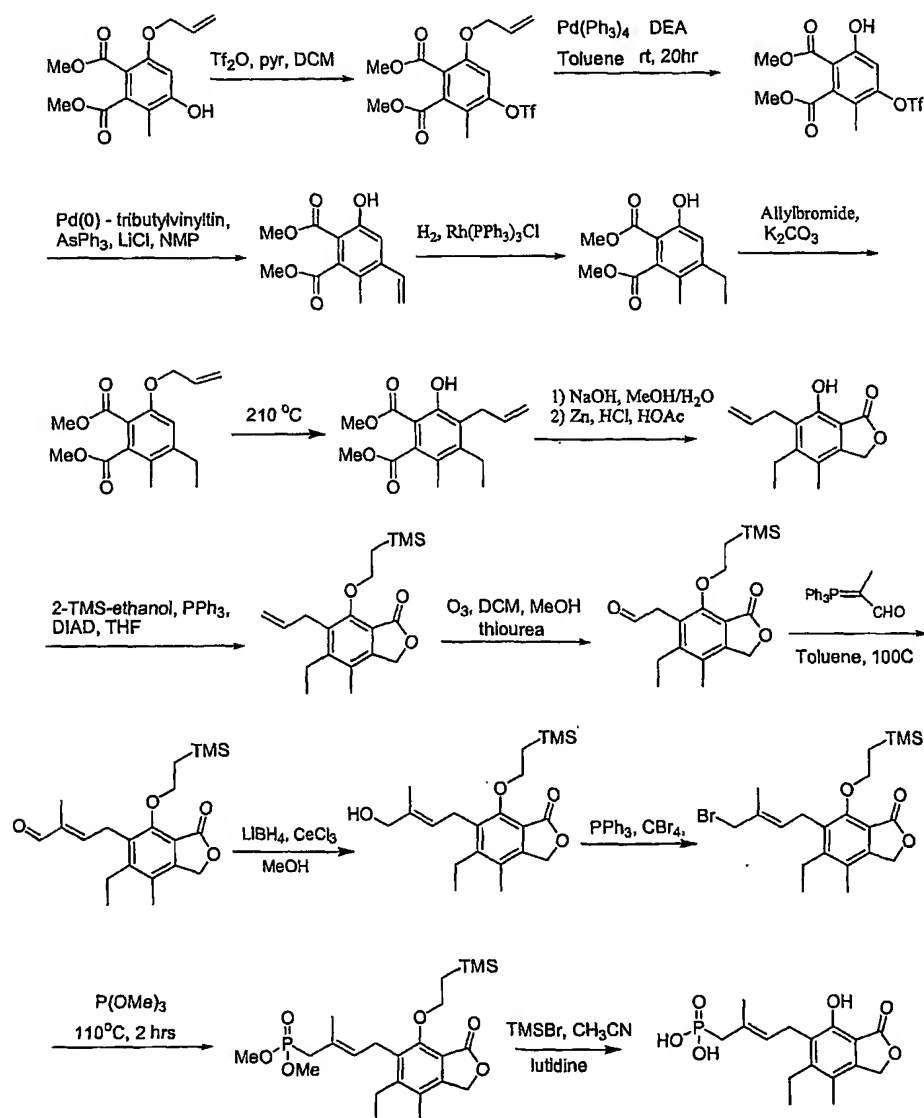
[2-Ethyl-4-(4-hydroxy-6-methoxy-7-methyl-3-oxo-1,3-dihydro-isobenzofuran-5-yl)-but-2-enyloxymethyl]-phosphonic acid

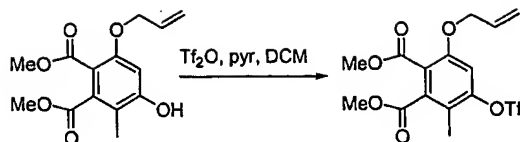
To a solution of {2-ethyl-4-[6-methoxy-7-methyl-3-oxo-4-(2-trimethylsilyl-ethoxy)-1,3-dihydro-isobenzofuran-5-yl]-but-2-enyloxymethyl}-phosphonic acid diisopropyl ester (347mg, 0.61mmol) in acetonitrile (5mL) was added 2,6-lutidine (0.71mL, 6.1mmol) and bromotrimethylsilane (0.786mL, 6.1mmol). The mixture was stirred at room temperature for 3 hours, quenched with methanol (5mL), concentrated, and partitioned between ethyl acetate and 1N HCl (aqueous). The organic layer was concentrated to give the free phosphonic acid as a colorless oil (205mg, 70%). This material (20mg) was dissolved in a solution of trifluoroacetic acid (0.3mL) and dichloromethane (2.7mL) and stirred for 30 minutes at ambient temperature. After concentration, the residue was purified by RP HPLC using a C18 column with a gradient of H₂O, 0.1% TFA-acetonitrile, 0.1% TFA to provide the

product, after lyophilization, as a white solid (10mg). ^1H NMR (300 MHz, CDCl_3) δ 1.007 (t, 3H, $J=7.5$ Hz), 2.13 (s, 3H), 2.32 (q, 2H, $J=7.5$ Hz), 3.41 (d, 2H, $J=6.3$ Hz), 3.56 (d, 2H, $J=9$ Hz), 3.75 (s, 3H), 3.95 (s, 2H), 5.16 (s, 2H), 5.43 (t, 1H, $J=6.3$ Hz) ppm; ^{31}P (121.4 MHz, CDCl_3) δ 22.8 ppm; MS (m/z) 385.2 $[\text{M}-\text{H}]^+$, 387.1 $[\text{M}+\text{H}]^+$.

Example 317

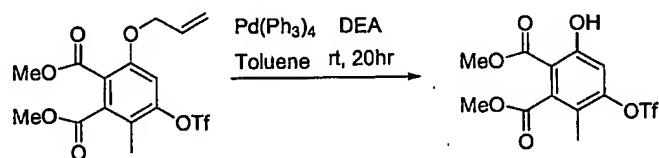
Representative compounds of the invention can be prepared as illustrated below.





6-Allyloxy-3-methyl-4-trifluoromethanesulfonyloxy-phthalic acid dimethyl ester

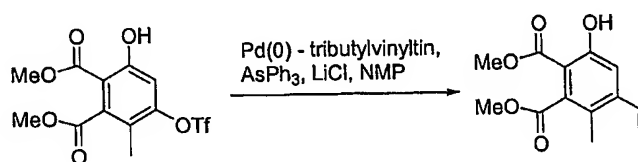
To a solution of 6-allyloxy-4-hydroxy-3-methyl-phthalic acid dimethyl ester (8.06 g, 28.8 mmol) [synthesized according to: J. W. Patterson, *Tetrahedron*, 1993, 49, 4789-4798] and pyridine (11.4 g, 144.0 mmol) in dichloromethane (DCM) (20 mL) at 0°C was added triflic anhydride (12.19 g, 43.2 mmol). The reaction was stirred at 0°C for 2 hours after which additional triflic anhydride (3 mL) was added. Stirring at 0°C was continued for an additional hour. The reaction mixture was poured into a mixture of DCM and HCl (1N). The layers were separated and the aqueous layer was extracted with DCM. The combined organic layers were dried over sodium sulfate. Filtration and evaporation of solvents *in vacuo* yielded a crude product, which was purified by silica gel chromatography to provide 8.39 g of the product as an oil. ^1H NMR (300 MHz, CDCl_3): δ = 2.32 (s, 3H), 3.89 (s, 6H), 4.60 (m, 2H), 5.33 (d, J = 9.3 Hz, 1H), 5.41 (d, J = 18.6 Hz, 1H), 5.95 (m, 1H), 6.95 (s, 1H) ppm; ^{19}F NMR (282 MHz, CDCl_3): δ = -74 ppm.



6-Hydroxy-3-methyl-4-trifluoromethanesulfonyloxy-phthalic acid dimethyl ester

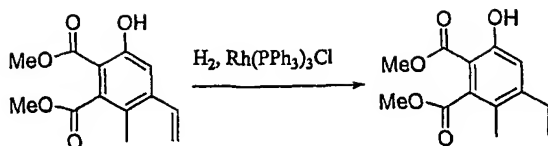
To a solution of 6-allyloxy-3-methyl-4-trifluoromethanesulfonyloxy-phthalic acid dimethyl ester (8.39 g, 20.3 mmol) in toluene (20 mL) was added tetrakis(triphenylphosphine) palladium (0.47 g, 0.40 mmol) and diethylamine (2.97 g, 40.86 mmol) at room temperature under an atmosphere of nitrogen.

Stirring at room temperature was continued until all starting material was consumed. The crude reaction mixture was partitioned between diethyl ether and HCl (0.1 N). The organic layer was washed with brine and dried over sodium sulfate. Filtration and evaporation of solvents *in vacuo* yielded a crude material, which was purified by silica gel chromatography to provide 4.16 g (55%) of the desired product as an off-white solid. ^1H NMR (300 MHz, CDCl_3): δ = 2.20 (s, 3H), 3.93 (s, 3H), 3.95 (s, 3H), 7.01 (s, 1H) ppm; ^{19}F NMR (282 MHz, CDCl_3): δ = -74 ppm.



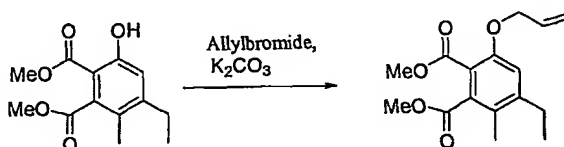
6-Hydroxy-3-methyl-4-vinyl-phthalic acid dimethyl ester

To a solution of 6-hydroxy-3-methyl-4-trifluoromethanesulfonyloxy-phthalic acid dimethyl ester (2.17 g, 5.85 mmol) in *N*-methyl pyrrolidinone (15 mL) was added lithium chloride (743 mg, 17.5 mmol) and triphenylarsine (179 mg, 0.585 mmol). Tributylvinyltin (2.04 g, 6.43 mmol) was added followed by tris(tribenzylideneacetone)dipalladium(0)-chloroform adduct (90 mg, 0.087 mmol). The reaction was placed under an atmosphere of nitrogen and heated at 60°C for 18 hours. The reaction was cooled to room temperature and poured onto a mixture of ice (20 g), EtOAc (40 mL), and potassium fluoride (1 g). Stirring was continued for 1 hour. The aqueous layer was extracted with EtOAc and the organic extracts filtered through Celite. The combined organic layers were washed with water and dried over sodium sulfate. Filtration and evaporation of solvents *in vacuo* yielded a crude material, which was purified by silica gel chromatography to provide 1.27 g (87%) of the product as an off-white solid. ^1H NMR (300 MHz, CDCl_3): δ = 2.16 (s, 3H), 3.91 (s, 3H), 3.92 (s, 3H), 5.46 (dd, J = 11.1, 1.2 Hz, 1H), 5.72 (dd, J = 17.1, 0.9 Hz, 1H), 6.86 (dd, J = 17.1, 11.1 Hz, 1H), 7.14 (s, 1H), 10.79 (s, 1H) ppm.



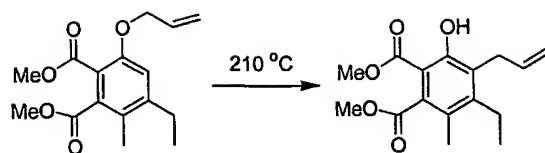
4-Ethyl-6-hydroxy-3-methyl-phthalic acid dimethyl ester

6-Hydroxy-3-methyl-4-vinyl-phthalic acid dimethyl ester (1.27 g, 5.11 mmol) was dissolved in benzene (10 mL) and EtOAc (10 mL). Tris(triphenylphosphine)rhodium chloride (150 mg) was added and the reaction was placed under an atmosphere of hydrogen. Stirring at room temperature was continued. After 14 hours, the solvents were removed *in vacuo* and the crude material was purified by silica gel chromatography to provide 1.14 g (88%) of the desired product as an off-white solid. ¹H NMR (300 MHz, CDCl₃): δ = 1.19 (t, *J* = 7.8 Hz, 3H), 2.10 (s, 3H), 2.60 (q, *J* = 7.8 Hz, 2H), 3.89 (s, 6H), 6.87 (s, 1H), 10.79 (s, 1H) ppm.



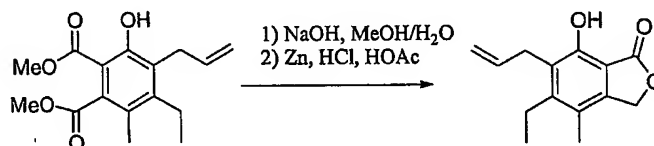
1 6-Allyloxy-4-ethyl-3-methyl-phthalic acid dimethyl ester

4-Ethyl-6-hydroxy-3-methyl-phthalic acid dimethyl ester (1.01 g, 4.02 mmol) was dissolved in DMF (5 mL). Potassium carbonate (3.33 g, 24.14 mmol) was added, followed by allylbromide (2.92 g, 24.14 mmol). The suspension was heated at 60°C. After 14 hours, the reaction was cooled to room temperature and filtered. The solvents were removed *in vacuo* and the crude material was purified by silica gel chromatography to provide 0.976 g (83%) of the desired product as a colorless oil. ¹H NMR (300 MHz, CDCl₃): δ = 1.16 (t, *J* = 7.2 Hz, 3H), 2.20 (s, 3H), 2.62 (q, *J* = 7.2 Hz, 2H), 3.83 (s, 3H), 3.84 (s, 3H), 4.57 (m, 2H), 5.26 (dd, *J* = 9.3, 1.5 Hz, 1H), 5.41 (dd, *J* = 13.5, 1.5 Hz, 1H), 5.98 (m, 1H), 6.82 (s, 1H) ppm.



4-Allyl-5-ethyl-3-hydroxy-6-methyl-phthalic acid dimethyl ester

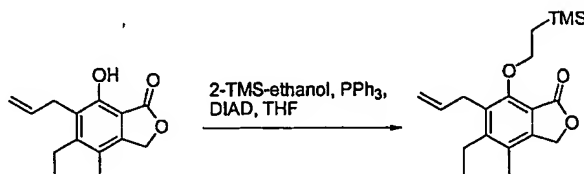
6-Allyloxy-4-ethyl-3-methyl-phthalic acid dimethyl ester (1.25 g, 4.28 mmol) was heated at 210°C under an atmosphere of nitrogen. After 14 hours, the reaction was cooled to room temperature. The crude material was purified by silica gel chromatography to provide 0.971 g (77%) of the desired product as a colorless oil. ¹H NMR (300 MHz, CDCl₃): δ = 1.14 (t, *J* = 7.8 Hz, 3H), 2.17 (s, 3H), 2.68 (q, *J* = 7.8 Hz, 2H), 3.49 (m, 2H), 3.86 (s, 3H), 3.89 (s, 3H), 4.89 – 5.01 (m, 2H), 5.93 (m, 1H), 11.22 (s, 1H) ppm.



5 6-Allyl-5-ethyl-7-hydroxy-4-methyl-3H-isobenzofuran-1-one

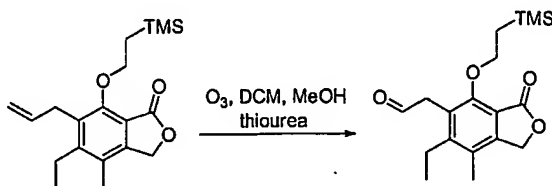
4-Allyl-5-ethyl-3-hydroxy-6-methyl-phthalic acid dimethyl ester (0.971 g, 3.32 mmol) was dissolved in MeOH (8 mL) at room temperature. A solution of sodium hydroxide (0.798 g, 19.95 mmol) in water (10 mL) was added and the suspension was heated at 55°C. After 16 hours, the reaction was cooled to room temperature and washed with diethyl ether. The aqueous layer was acidified (1N HCl) and the suspension was extracted with EtOAc. The combined organic layers were dried over sodium sulfate. Filtration and evaporation of solvents *in vacuo* yielded the desired bis acid as a white solid (0.846 g, 98%, *M*⁺ = 263). The bis acid was dissolved in acetic acid (6 mL) and HCl (conc., 1.5 mL). The reaction was heated at 80°C. Zn dust (0.635 g, 9.72 mmol, each) was added in portions every hour for 7 hours. Stirring at 80°C was continued for additional 10 hours. The reaction was cooled to room temperature, and water was added. The resultant suspension was extracted with EtOAc. The combined organic extracts were washed with sodium bicarbonate solution and dried over sodium sulfate.

Filtration and evaporation of solvents *in vacuo* yielded the crude product, which was purified by silica gel chromatography to provide 0.375 g (50%) of the product as a white solid. ^1H NMR (300 MHz, CDCl_3): δ = 1.14 (t, J = 7.5 Hz, 3H), 2.18 (s, 3H), 2.71 (q, J = 7.5 Hz, 2H), 3.49 (m, 2H), 4.95 (d, J = 17.1 Hz, 1H), 5.02 (d, J = 10.2 Hz, 1H), 5.23 (s, 2H), 5.98 (m, 1H), 7.66 (s, 1H) ppm.



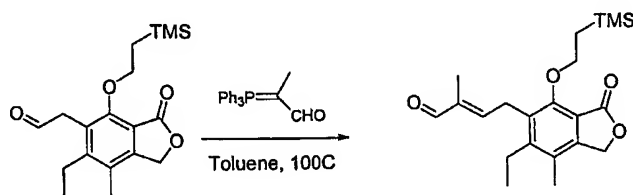
5-6-Allyl-5-ethyl-4-methyl-7-(2-trimethylsilyl-ethoxy)-3H-isobenzofuran-1-one

To a solution of 6-allyl-5-ethyl-7-hydroxy-4-methyl-3H-isobenzofuran-1-one (199 mg, 0.857 mmol), PPh_3 (337 mg, 1.286 mmol), and 2-trimethylsilyl-ethanol in THF (3 mL) at 0°C was added diisopropyl azodicarboxylate (259 mg, 1.286 mmol). The resulting yellow solution was allowed to warm to room temperature and stirred for one hour. The solvent was removed *in vacuo* and the crude material was dissolved in diethyl ether (3 mL). Hexanes (1.5 mL) were added. Triphenylphosphine oxide was removed by filtration and the filtrate was concentrated and purified by silica gel chromatography to provide the desired product (261 mg, 92 %) as a clear oil. ^1H NMR (300 MHz, CDCl_3): δ = 0.04 (s, 9H), 1.15 (t, J = 7.8 Hz, 3H), 1.25 (m, 2H), 2.20 (s, 3H), 2.73 (q, J = 7.8 Hz, 2H), 3.54 (m, 2H), 4.28 (m, 2H), 4.95 (d, J = 17.1 Hz, 1H), 5.02 (d, J = 10.2 Hz, 1H), 5.15 (s, 2H), 5.95 (m, 1H) ppm.



[6-Ethyl-7-methyl-3-oxo-4-(2-trimethylsilyl-ethoxy)-1,3-dihydroisobenzofuran-5-yl]-acetaldehyde

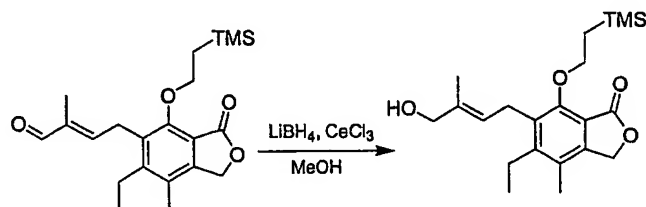
A solution of 6-allyl-5-ethyl-4-methyl-7-(2-trimethylsilyl-ethoxy)-3H-isobenzofuran-1-one (261 mg, 0.788 mmol) in MeOH (5 mL), CH₂Cl₂ (5 mL) and pyridine (50 μ L) was cooled to -78°C using a dry ice/acetone bath according to the procedure of Smith, D. B. *et al.*, *J. Org. Chem.*, 1996, 61, 6, 2236. A stream of ozone was bubbled through the reaction *via* a gas dispersion tube until the reaction became blue in color (15 minutes). The ozone line was replaced with a stream of nitrogen and bubbling continued for another 15 minutes, by which time the blue color had disappeared. To this solution, at -78°C , was added thiourea (59.9 mg, 0.788 mmol) in one portion, and the cooling bath was removed. The reaction was allowed to warm to room temperature and stirred for 15 hours. The reaction mixture was filtered and then partitioned between CH₂Cl₂ and water. The aqueous layer was extracted with CH₂Cl₂ one more time and the organic extracts were combined, washed with aqueous 1N HCl, saturated NaHCO₃ and brine and dried over sodium sulfate. Filtration and evaporation of solvents in vacuo yielded the crude product, which was purified by silica gel chromatography to afford 181 mg (69 %) of the product as a white solid. ¹H NMR (300 MHz, CDCl₃): δ = 0.04 (s, 9H), 1.11 (t, J = 7.5 Hz, 3H), 1.19 (m, 2H), 2.21 (s, 3H), 2.66 (q, J = 7.5 Hz, 2H), 3.90 (s, 2H), 4.36 (m, 2H), 5.18 (s, 2H), 9.71 (s, 1H) ppm.



4-[6-Ethyl-7-methyl-3-oxo-4-(2-trimethylsilyl-ethoxy)-1,3-dihydroisobenzofuran-5-yl]-2-methyl-but-2-enal

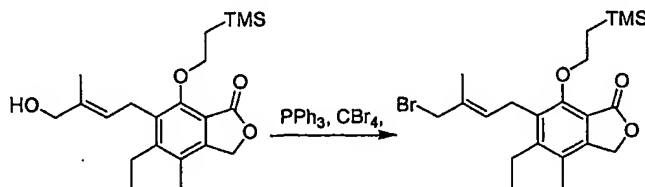
[6-Ethyl-7-methyl-3-oxo-4-(2-trimethylsilyl-ethoxy)-1,3-dihydroisobenzofuran-5-yl]-acetaldehyde (90 mg, 0.269 mmol) and 2-(triphenylphosphorylidene)-propionaldehyde (72.9 mg, 0.23 mmol) in toluene (3 mL) were heated at 100°C . After 15 hours, a second portion of 2-(triphenylphosphanylidene)-propionaldehyde (33 mg, 0.11 mmol) was added and the reaction mixture was heated for additional 9 hours. The toluene was removed *in*

vacuo, and the residue was purified by silica gel chromatography to provide 77.6 mg (77%) of the desired product as a pale yellow oil. ^1H NMR (300 MHz, CDCl_3): δ = 0.03 (s, 9H), 1.15 (t, J = 7.5 Hz, 3H), 1.21 (m, 2H), 1.93 (s, 3H), 2.21 (s, 3H), 2.71 (q, J = 7.5 Hz, 2H), 3.82 (d, J = 6.9 Hz, 2H), 4.34 (m, 2H), 5.18 (s, 2H), 6.38 (m, 1H), 9.35 (s, 1H) ppm.



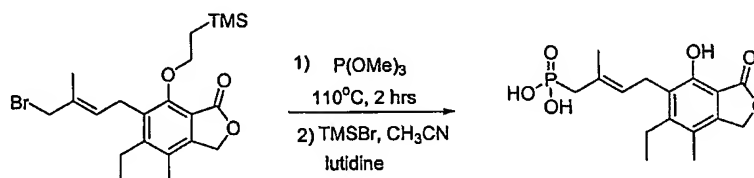
5-Ethyl-6-(4-hydroxy-3-methyl-but-2-enyl)-4-methyl-7-(2-trimethylsilyl-ethoxy)-3H-isobenzofuran-1-one

4-[6-Ethyl-7-methyl-3-oxo-4-(2-trimethylsilyl-ethoxy)-1,3-dihydroisobenzofuran-5-yl]-2-methyl-but-2-enal (77.6 mg, 0.207 mmol) was dissolved in MeOH (4 mL). A solution of CeCl_3 (51.1 mg, 0.207 mmol) in MeOH/water (9/1, 0.66 mL) was added and the solution was cooled to 0°C . A solution of lithium borohydride in THF (2M, 0.105 mL) was added dropwise. After 15 minutes, the reaction was quenched with 1N HCl (0.5 mL). The MeOH was removed *in vacuo* and the crude material was partitioned between DCM and water. The aqueous layer was extracted with DCM and the combined organic layers were washed with sodium bicarbonate solution and dried over sodium sulfate. Filtration and evaporation of solvents yielded a crude oil, which was purified by silica gel chromatography to provide 57.2 mg (73%) of the desired product. ^1H NMR (300 MHz, CDCl_3): δ = 0.04 (s, 9H), 1.15 (t, J = 7.8 Hz, 3H), 1.26 (m, 2H), 1.86 (s, 3H), 2.19 (s, 3H), 2.72 (q, J = 7.8 Hz, 2H), 3.52 (d, J = 6.3 Hz, 2H), 3.99 (s, 2H), 4.34 (m, 2H), 5.14 (s, 2H), 5.32 (m, 1H) ppm.



6-(4-Bromo-3-methyl-but-2-enyl)-5-ethyl-4-methyl-7-(2-trimethylsilylanyl-ethoxy)-3H-isobenzofuran-1-one

5-Ethyl-6-(4-hydroxy-3-methyl-but-2-enyl)-4-methyl-7-(2-trimethylsilylanyl-ethoxy)-3H-isobenzofuran-1-one (57.2 mg, 0.152 mmol) was dissolved in DCM (3.5 mL). Polymer-bound triphenylphosphine (3 mmol/g, 152.1 mg) was added and the mixture was mechanically stirred at room temperature. Carbon tetrabromide (151.3 mg, 0.456 mmol) was added and the solution was stirred at room temperature. After 2 hours, the reaction was filtered and the solvent was removed *in vacuo*. The crude material was purified by silica gel chromatography to provide 58.0 mg (87%) of the desired product. ¹H NMR (300 MHz, CDCl₃): δ = 0.04 (s, 9H), 1.15 (t, *J* = 7.8 Hz, 3H), 1.25 (m, 2H), 1.95 (s, 3H), 2.20 (s, 3H), 2.70 (q, *J* = 7.8 Hz, 2H), 3.52 (d, *J* = 6.3 Hz, 2H), 3.94 (s, 2H), 4.28 (m, 2H), 5.14 (s, 2H), 5.50 (m, 1H) ppm.



{4-[6-Ethyl-7-methyl-3-oxo-4-(2-trimethylsilylanyl-ethoxy)-1,3-dihydro-isobenzofuran-5-yl]-2-methyl-but-2-enyl}-phosphonic acid

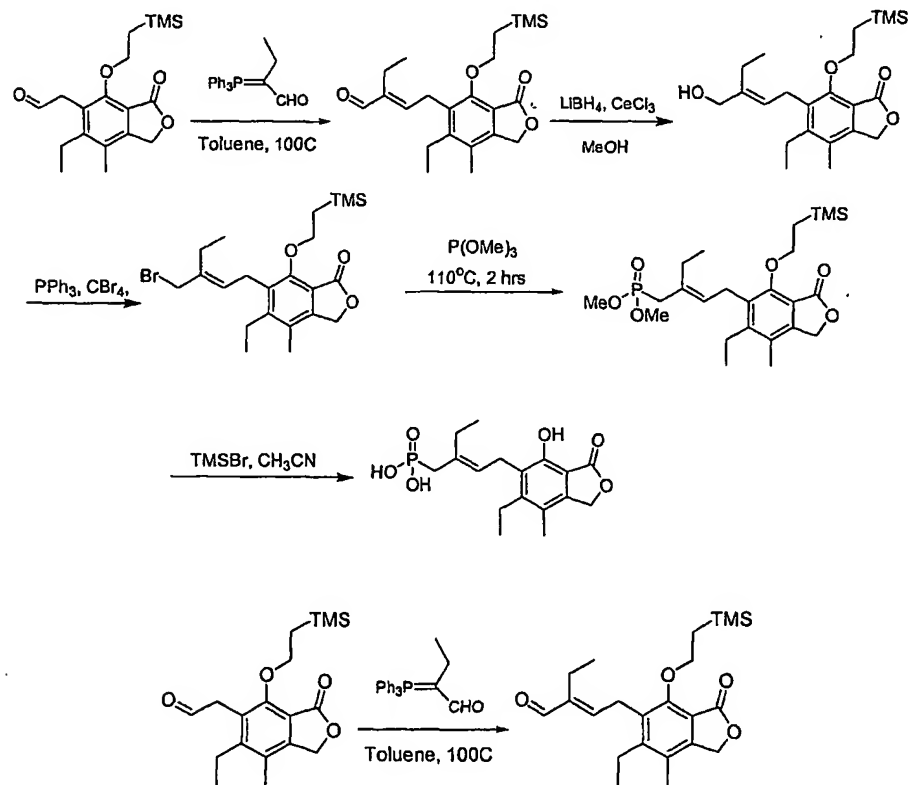
A solution of 4-[6'-ethyl-7'-methyl-3'-oxo-4'-(2''-trimethylsilylanyl-ethoxy)-1',3'-dihydro-isobenzofuran-5'-yl]-2-methyl-but-2-enyl bromide (58 mg, 0.132 mmol) in trimethylphosphite (0.8 mL) was heated at 110°C. After 2 hours the reaction was complete. The reaction was cooled to room temperature and the excess trimethylphosphite was removed *in vacuo*. The crude material was used in the next step without further purification.

The crude product of the Arbuzov reaction was dissolved in MeCN (0.8 mL). Trimethylsilyl bromide (202.2 mg, 1.321 mmol) was added and the reaction was stirred at room temperature. After 15 minutes, lutidine (155.7 mg, 1.453 mmol) was added and stirring at room temperature was continued. After 2 hours, additional trimethylsilyl bromide (202.2 mg, 1.321 mmol) was added and stirring at room temperature was continued. After 4 hours, the reaction was

quenched with MeOH (2 mL). The solvents were evaporated *in vacuo*, and the crude material was purified by RP-HPLC (eluent: water / MeCN). The product-containing fractions were combined and lyophilized to yield 2.3 mg (5.1%) of the free phosphonic acid. ^1H NMR (300 MHz, DMSO- d_6): δ = 1.07 (t, J = 7.5 Hz, 3H), 1.84 (s, 3H), 2.14 (s, 3H), 2.64 (q, J = 7.5 Hz, 2H), 3.34 (m, 4H), 5.06 (m, 1H), 5.25 (s, 2H) ppm; ^{31}P NMR (121 MHz, DMSO- d_6): δ = 22.19 ppm; MS = 341 [M^+ +1].

Example 318

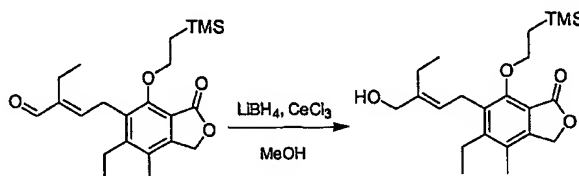
Representative compounds of the invention can be prepared as illustrated below.



[2-Ethyl-4-[6-ethyl-7-methyl-3-oxo-4-(2-trimethylsilyl-ethoxy)-1,3-dihydro-isobenzofuran-5-yl]-but-2-enal

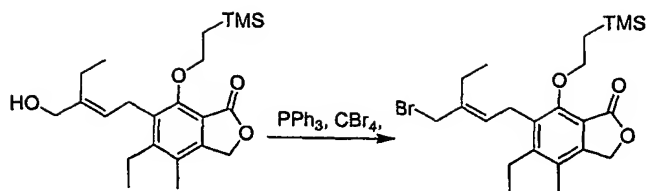
[6-Ethyl-7-methyl-3-oxo-4-(2-trimethylsilyl-ethoxy)-1,3-dihydro-isobenzofuran-5-yl]-acetaldehyde (90 mg, 0.269 mmol) and 2-(triphenyl-

phosphorylidene)-butyraldehyde (98.4 mg, 0.296 mmol) in toluene (3 mL) were heated at 100°C. After 15 hours, a second portion of 2-(triphenylphosphanylidene)-butyraldehyde (98.4 mg, 0.296 mmol) was added and the reaction mixture was heated for additional 33 hours. After concentration, the residue was purified by silica gel chromatography to provide 50.3 mg (48%) of the desired product as a pale yellow oil.



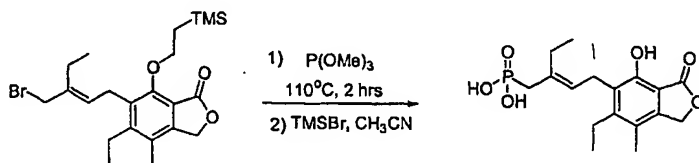
5-Ethyl-6-(3-hydroxymethyl-pent-2-enyl)-4-methyl-7-(2-trimethylsilyl-ethoxy)-3H-isobenzofuran-1-one

2-Ethyl-4-[6-ethyl-7-methyl-3-oxo-4-(2-trimethylsilyl-ethoxy)-1,3-dihydro-isobenzofuran-5-yl]-but-2-enal (50.3 mg, 0.129 mmol) was dissolved in MeOH (3 mL). A solution of CeCl_3 (31.9 mg, 0.129 mmol) in MeOH/water (9/1, 0.66 mL) was added and the solution was cooled to 0°C. A solution of lithium borohydride in THF (2M, 0.065 mL) was added dropwise. After 10 minutes, the reaction was quenched with 1N HCl (0.5 mL). The methanol was removed *in vacuo* and the crude material was partitioned between DCM and water. The aqueous layer was extracted with DCM and the combined organic layers were washed with sodium bicarbonate solution and were dried over sodium sulfate. Filtration and evaporation of solvents *in vacuo* yielded a crude oil, which was purified by silica gel chromatography to provide 35.4 mg (70%) of the desired product. ^1H NMR (300 MHz, CDCl_3): δ = 0.04 (s, 9H), 1.10 – 1.19 (m, 6H), 1.26 (m, 2H), 2.19 (s, 3H), 2.32 (q, J = 7.5 Hz, 2H), 2.72 (q, J = 7.5 Hz, 2H), 3.54 (d, J = 6.6 Hz, 2H), 4.05 (s, 2H), 4.26 (m, 2H), 5.14 (s, 2H), 5.27 (m, 1H) ppm.



6-(3-Bromomethyl-pent-2-enyl)-5-ethyl-4-methyl-7-(2-trimethylsilyl-ethoxy)-3H-isobenzofuran-1-one

5-Ethyl-6-(3-hydroxymethyl-pent-2-enyl)-4-methyl-7-(2-trimethylsilyl-ethoxy)-3H-isobenzofuran-1-one (35.4 mg, 0.090 mmol) was dissolved in DCM (3.0 mL). Polymer-bound triphenylphosphine (3 mmol/g, 90.7 mg) was added, and the mixture was mechanically stirred at room temperature. Carbon tetrabromide (90.2 mg, 0.272 mmol) was added and the solution was stirred at room temperature. After 2 hours, the reaction was filtered and the solvent was removed *in vacuo*. The crude material was purified by silica gel chromatography to provide 32.0 mg (78%) of the desired product. The material was used in the next step without further characterization.



[2-Ethyl-4-(6-ethyl-4-hydroxy-7-methyl-3-oxo-1,3-dihydro-isobenzofuran-5-yl)-but-2-enyl]-phosphonic acid

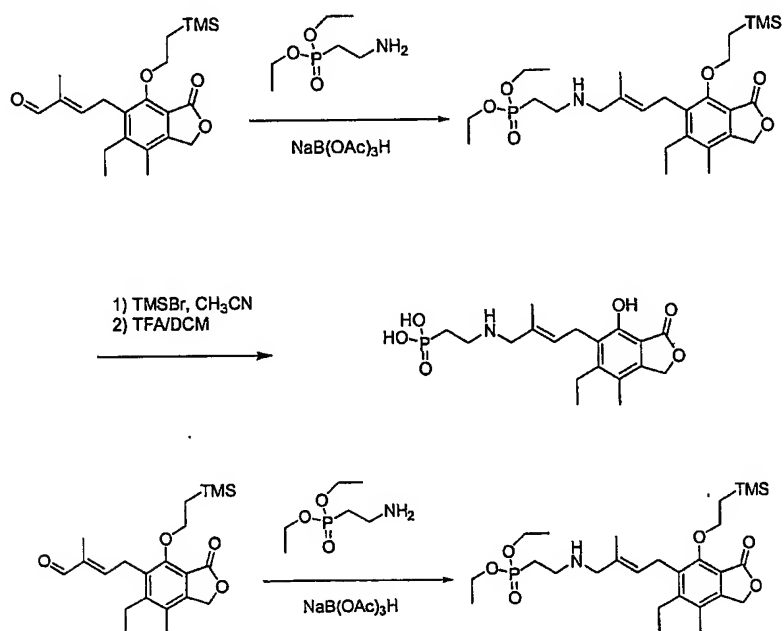
A solution of 6-(3-bromomethyl-pent-2-enyl)-5-ethyl-4-methyl-7-(2-trimethylsilyl-ethoxy)-3H-isobenzofuran-1-one (32 mg, 0.070 mmol) in trimethylphosphite (0.8 mL) was heated at 110 °C. After 2 hours, the reaction was complete. The reaction was cooled to room temperature and the excess trimethylphosphite was removed *in vacuo*. The crude material was used in the next step without further purification.

The crude product of the Arbuzov reaction was dissolved in MeCN (0.8 mL). Trimethylsilyl bromide (108.0 mg, 0.706 mmol) was added and the reaction was stirred at room temperature. After 2 hours, a second batch of

trimethylsilyl bromide (108.0 mg, 0.706 mmol) was added. After 3 hours, the reaction was quenched with MeOH (2 mL). The solvents were evaporated *in vacuo* and the crude material was purified by RP-HPLC (eluent: water / MeCN). The product-containing fractions were combined and lyophilized to yield 15.7 mg (63%) of the product. ^1H NMR (300 MHz, DMSO- d_6): δ = 0.98 - 1.09 (m, 6H), 2.10 (s, 3H), 2.30 (m, 2H), 2.64 (q, J = 7.5 Hz, 2H), 3.38 (m, 4H), 5.03 (m, 1H), 5.25 (s, 2H) ppm; ^{31}P NMR (121 MHz, DMSO- d_6): δ = 22.26 ppm; MS = 355 [M^+ +1].

Example 319

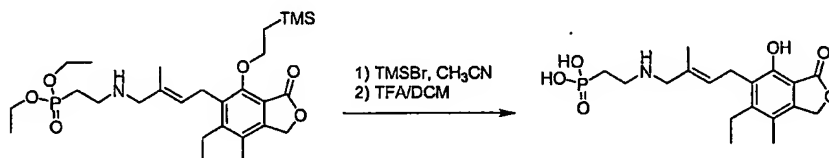
Representative compounds of the invention can be prepared as illustrated below.



(2-{4-[6-Ethyl-7-methyl-3-oxo-4-(2-trimethylsilylanyl-ethoxy)-1,3-dihydro-isobenzofuran-5-yl]-2-methyl-but-2-enylamino}-ethyl)-phosphonic acid diethyl ester

4-[6-Ethyl-7-methyl-3-oxo-4-(2-trimethylsilylanyl-ethoxy)-1,3-dihydro-isobenzofuran-5-yl]-2-methyl-but-2-enal (19.7 mg, 0.052 mmol) and aminoethylphosphonic acid diethylester oxalate salt (15.6 mg, 0.057 mmol) were

dissolved in DMF (0.5 mL). Acetic acid (15.7 mg, 0.263 mmol) was added, followed by sodium triacetoxyborohydride (22.3 mg, 0.105 mmol). After 4 hours, the crude reaction mixture was purified by RP-HPLC (eluent: water/MeCN) to provide 27.7 mg (97%) of the desired product after lyophilization. ^1H NMR (300 MHz, CDCl_3): δ = 0.04 (s, 9H), 1.14 (t, J = 7.5 Hz, 3H), 1.26 (m, 2H), 1.30 (t, J = 7.2 Hz, 6H), 1.95 (s, 3H), 2.19 (s, 3H), 2.23 (m, 2H), 2.68 (q, J = 7.5 Hz, 2H), 3.18 (m, 2H), 3.53 (s, 2H), 4.13 (m, 4H), 4.28 (m, 2H), 5.15 (s, 2H), 5.51 (m, 1H) ppm; ^{31}P NMR (121 MHz, CDCl_3): δ = 27.39 ppm; MS = 540 [M^+ +1].

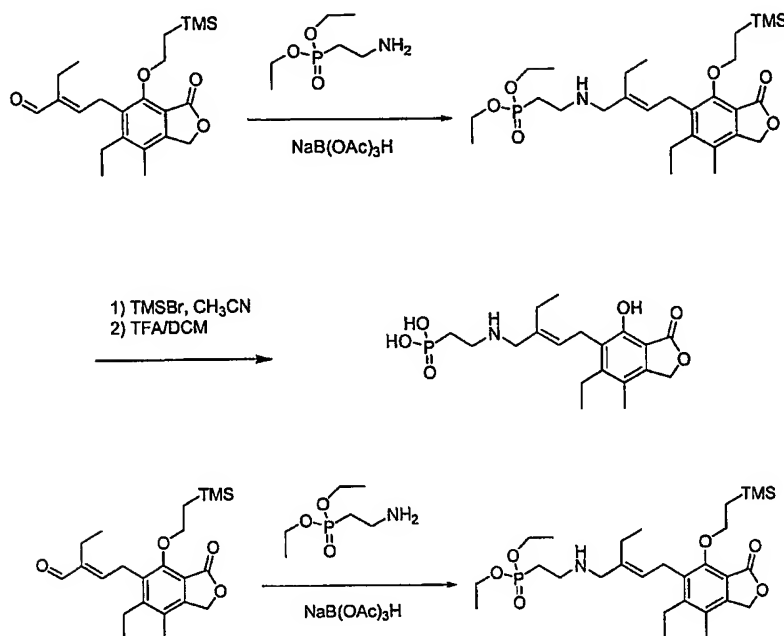


{2-[4-(6-Ethyl-4-hydroxy-7-methyl-3-oxo-1,3-dihydro-isobenzofuran-5-yl)-2-methyl-but-2-enylamino]-ethyl}-phosphonic acid:

(2-{4-[6-Ethyl-7-methyl-3-oxo-4-(2-trimethylsilyl-ethoxy)-1,3-dihydro-isobenzofuran-5-yl]-2-methyl-but-2-enylamino}-ethyl)-phosphonic acid diethyl ester (27.7 mg, 0.051 mmol) was dissolved in DMF (0.5 mL) and DCM (0.5 mL). Trimethylsilyl bromide (78.3 mg, 0.512 mmol) was added and the reaction was stirred at room temperature. After 20 hours, the reaction was quenched with MeOH (0.3 mL). The solvents were evaporated *in vacuo* and the crude material was purified by RP-HPLC (eluent: water/MeCN). The product-containing fractions were combined and lyophilized to yield 14.2 mg (57%) of the free phosphonic acid [MS: 484 M^+ +1].

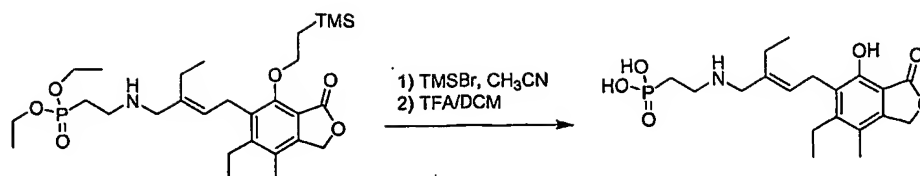
The material was dissolved in DCM (0.5 mL). TFA (0.05 mL) was added and stirring at room temperature was continued. After 20 minutes, the solvents were removed *in vacuo* and the crude material was purified by RP-HPLC (eluent: water/MeCN * 0.1% TFA). The product-containing fractions were combined and lyophilized to yield 7.6 mg (52%) of the product as the TFA salt. ^1H NMR (300 MHz, $\text{DMSO}-d_6$): δ = 1.07 (t, J = 7.5 Hz, 3H), 1.84 (s, 3H), 1.90 (m, 2H), 2.11 (s, 3H), 2.63 (q, J = 7.5 Hz, 2H), 2.99 (m, 2H), 3.43 (d, J =

6.3 Hz, 2H), 3.51 (s, 2H), 5.26 (s, 2H), 5.45 (m, 1H) ppm; ^{31}P NMR (121 MHz, DMSO- d_6): δ = 20.02 ppm; MS = 384 [M^+ +1].



(2-{2-Ethyl-4-[6-ethyl-7-methyl-3-oxo-4-(2-trimethylsilanyl-ethoxy)-1,3-dihydro-isobenzofuran-5-yl]-but-2-enylamino}-ethyl)-phosphonic acid diethyl ester

2-Ethyl-4-[6-ethyl-7-methyl-3-oxo-4-(2-trimethylsilanyl-ethoxy)-1,3-dihydro-isobenzofuran-5-yl]-but-2-enal (26.6 mg, 0.068 mmol) and aminoethylphosphonic acid diethylester oxalate salt (20.4 mg, 0.075 mmol) were dissolved in DMF (0.8 mL). Acetic acid (20.5 mg, 0.342 mmol) was added, followed by sodium triacetoxyborohydride (27.6 mg, 0.137 mmol). After 8 hours, the crude reaction mixture was purified by RP-HPLC (eluent: water/MeCN) to provide 24.9 mg (65%) of the desired product after lyophilization. ^1H NMR (300 MHz, CDCl_3): δ = 0.05 (s, 9H), 1.10 - 1.24 (m, 8H), 1.35 (t, J = 7.5 Hz, 6H), 2.19 (s, 3H), 2.23 (m, 2H), 2.35 (q, J = 7.8 Hz, 2H), 2.70 (q, J = 7.2 Hz, 2H), 3.25 (m, 2H), 3.56 (m, 4H), 4.15 (m, 4H), 4.29 (m, 2H), 5.15 (s, 2H), 5.47 (m, 1H) ppm; ^{31}P NMR (121 MHz, CDCl_3): δ = 27.71 ppm; MS = 554 [M^+ +1].



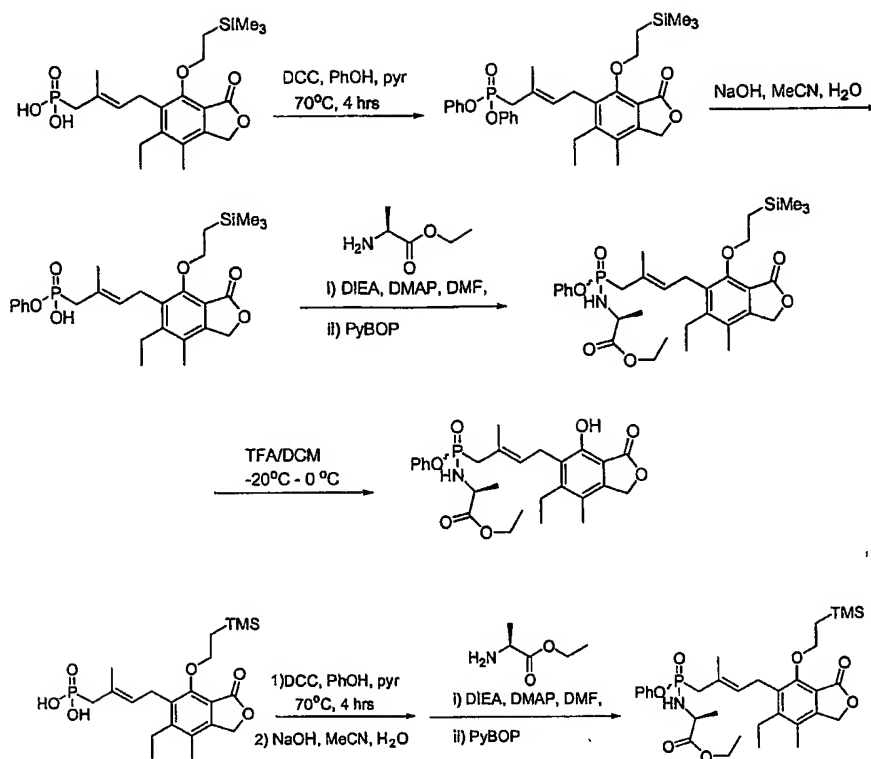
{2-[2-Ethyl-4-(6-ethyl-4-hydroxy-7-methyl-3-oxo-1,3-dihydro-isobenzofuran-5-yl)-but-2-enylamino]-ethyl}-phosphonic acid

(2-{2-Ethyl-4-[6-ethyl-7-methyl-3-oxo-4-(2-trimethylsilyl-ethoxy)-1,3-dihydro-isobenzofuran-5-yl]-but-2-enylamino}-ethyl)-phosphonic acid diethyl ester (24.9 mg, 0.045 mmol) was dissolved in DMF (0.5 mL) and DCM (0.5 mL). Trimethylsilyl bromide (68.7 mg, 0.449 mmol) was added and the reaction was stirred at room temperature. After 20 hours, the reaction was quenched with MeOH (0.15 mL). The solvents were evaporated *in vacuo* and the crude material was purified by RP-HPLC (eluent: water/MeCN). The product-containing fractions were combined and lyophilized to yield 8.0 mg of the free phosphonic acid [MS: 498 $M^+ + 1$].

This material was dissolved in DCM (0.5 mL). TFA (0.05 mL) was added, and stirring at room temperature was continued. After 20 minutes, the solvents were removed *in vacuo* and the crude material was purified by RP-HPLC (eluent: water/MeCN * 0.1% TFA). The product-containing fractions were combined and lyophilized to yield 4.4 mg (54%) of the product as the TFA salt. ^1H NMR (300 MHz, DMSO- d_6): δ = 1.05 (m, 6H), 1.60 (m, 2H), 2.10 (s, 3H), 2.67 (q, J = 7.5 Hz, 2H), 2.63 (q, J = 6.9 Hz, 2H), 2.93 (m, 2H), 3.45 (m, 4H), 5.24 (s, 2H), 5.36 (m, 1H) ppm.; ^{31}P NMR (121 MHz, DMSO- d_6): δ = 16.93 ppm; MS = 398 [$M^+ + 1$].

Example 320

Representative compounds of the invention can be prepared as illustrated below.



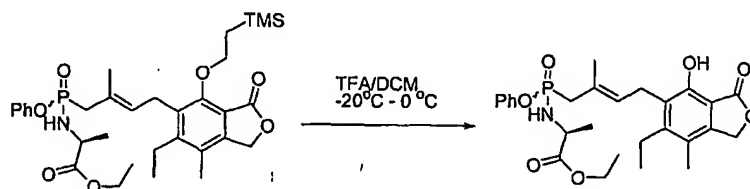
2-({4-[6-Ethyl-7-methyl-3-oxo-4-(2-trimethylsilyl-ethoxy)-1,3-dihydro-isobenzofuran-5-yl]-2-methyl-but-2-enyl}-phenoxy-phosphinoylamino)-propionic acid ethyl ester

4-[6'-ethyl-7'-methyl-3'-oxo-4'-(2''-trimethylsilyl-ethoxy)-1',3'-dihydro-isobenzofuran-5'-yl]-2-methyl-but-2-en-phosphonic acid (44.8 mg, 0.101 mmol), dicyclohexylcarbodiimide (52.6 mg, 0.254 mmol), and phenol (95.8 mg, 1.018 mmol) were dissolved in pyridine (0.3 mL) and heated at 70°C for 4 hours. The reaction mixture was cooled to room temperature and the pyridine was removed *in vacuo*. The crude material was partitioned between DCM and HCl (0.1N). The aqueous layer was extracted with DCM and the combined organic layers were dried over sodium sulfate. Filtration and evaporation of solvents *in vacuo* yielded a crude material, which was used in the next step without further purification.

The crude material was dissolved in MeCN (0.8 mL) and water (0.3 mL). Aqueous sodium hydroxide solution (2N, 0.8 mL) was added in portions (0.2 mL). After all starting material was consumed, the organic solvent was removed *in vacuo* and the crude material was partitioned between chloroform and aqueous

HCl (1N). The aqueous layer was extracted with chloroform. The combined organic layers were dried over sodium sulfate. Filtration and evaporation of solvents yielded the crude product as a mixture of mono phenyl ester and the symmetrical anhydride.

The crude material of the previous step and ethyl (*L*)-alanine hydrochloride salt (78.1 mg, 0.509 mmol) were dissolved in DMF (0.4 mL). DMAP (1.2 mg, catalytic) was added, followed by diisopropylethylamine (131.3 mg, 1.018 mmol). Stirring at room temperature was continued. After 20 minutes, complete conversion of the anhydride was observed. After 2 hours, PyBOP (101 mg, 0.202 mmol) was added and stirring at room temperature was continued. The reaction was filtered and the crude reaction solution was purified by RP-HPLC (eluent: water/MeCN). The product-containing fractions were combined and lyophilized to yield the product (15.7 mg, 25% over three steps) as a white powder. ^1H NMR (300 MHz, CDCl_3): δ = 0.03 (s, 9H), 1.13 – 1.28 (m, 8H), 2.03 (s, 3H), 2.19 (s, 3H), 2.62 – 2.74 (m, 4H), 3.38 (m, 1H), 3.53 (t, J = 6.3 Hz, 2H), 4.03 (m, 3H), 4.30 (m, 2H), 5.14 (s, 2H), 5.31 (m, 1H), 7.11 – 7.17 (m, 3H), 7.25 – 7.30 (m, 2H) ppm; ^{31}P NMR (121 MHz, CDCl_3): δ = 27.04, 27.73 ppm; MS = 615 [$\text{M}^+ + 1$].



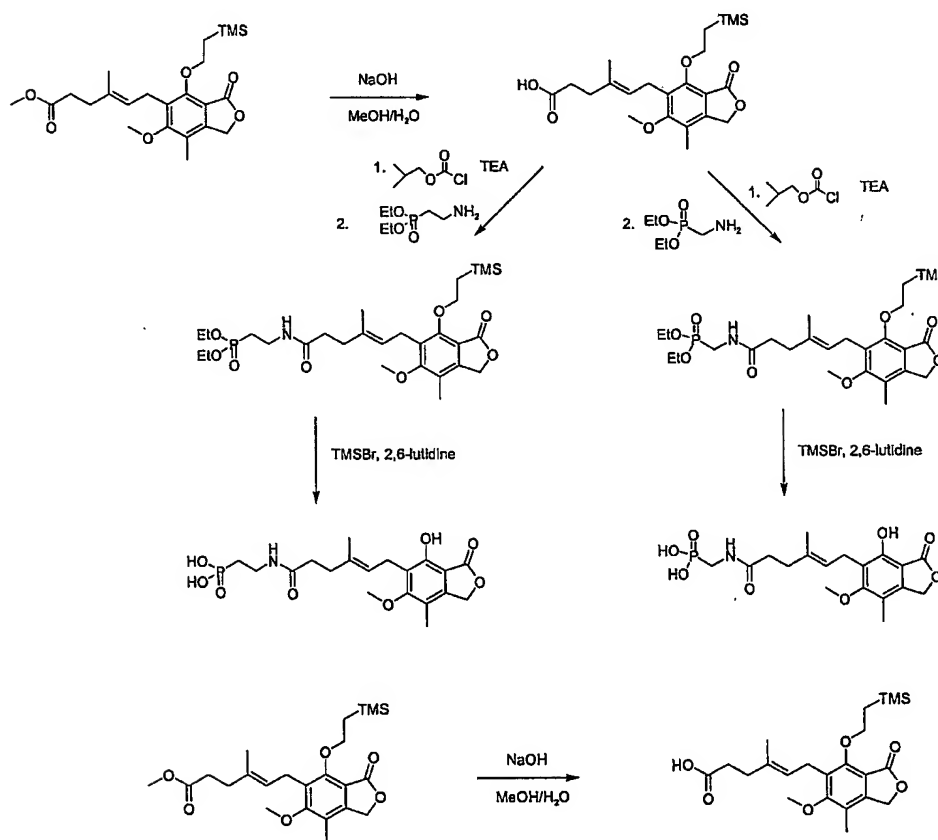
2-[[4-(6-Ethyl-4-hydroxy-7-methyl-3-oxo-1,3-dihydro-isobenzofuran-5-yl)-2-methyl-but-2-enyl]-phenoxy-phosphinoylamino}-propionic acid ethyl ester

2-({4-[6-Ethyl-7-methyl-3-oxo-4-(2-trimethylsilylanyl-ethoxy)-1,3-dihydro-isobenzofuran-5-yl]-2-methyl-but-2-enyl}-phenoxy-phosphinoylamino)-propionic acid ethyl ester (7.5 mg, 0.012 mmol) was dissolved in TFA/DCM (10%, 0.3 mL) at -20°C . The reaction mixture was warmed to 0°C and stirred at this temperature for 45 minutes. Pyridine (0.09 mL) was added the solvents were removed *in vacuo*. The crude material was

purified by RP-HPLC (eluent: water/MeCN). The product-containing fractions were combined and lyophilized, yielding a white powder (5.5 mg, 87%). ^1H NMR (300 MHz, CDCl_3): δ = 1.12 – 1.29 (m, 6H), 2.03 (s, 3H), 2.17 (s, 3H), 2.65 – 2.74 (m, 4H), 3.38 (m, 1H), 3.53 (t, J = 6.3 Hz, 2H), 4.03 (m, 3H), 5.22 (s, 2H), 5.36 (m, 1H), 7.11 – 7.16 (m, 3H), 7.24 – 7.30 (m, 2H), 7.72 (m, 1H) ppm; ^{31}P NMR (121 MHz, CDCl_3): δ = 27.11, 27.57 ppm; MS = 515 [M^+ +1].

Example 321

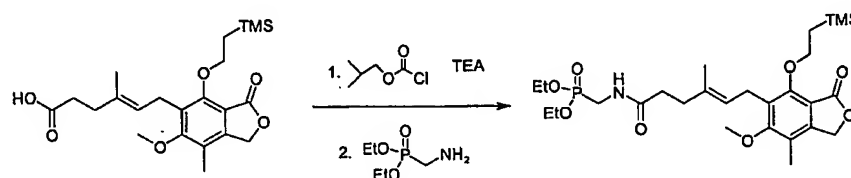
Representative compounds of the invention can be prepared as illustrated below.



6-[6-Methoxy-7-methyl-3-oxo-4-(2-trimethylsilyl-ethoxy)-1,3-dihydro-isobenzofuran-5-yl]-4-methyl-hex-4-enoic acid

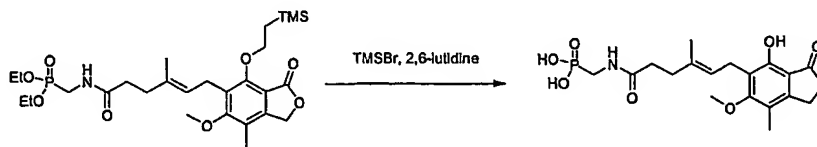
A mixture of 6-[6-methoxy-7-methyl-3-oxo-4-(2-trimethylsilyl-ethoxy)-1,3-dihydro-isobenzofuran-5-yl]-4-methyl-hex-4-enoic acid methyl ester

(1.5 g, 3.45 mmol) and sodium hydroxide (552 mg) in a mixture of methanol (20 mL) and water (7 mL) was stirred at room temperature for one hour. The solution was acidified with 1N HCl. The precipitate was collected by suction filtration and washed with water to give the desired product (1.2g, 83%). ^1H NMR (300 MHz, CDCl_3) δ 0.02 (s, 9H), 1.15- 1.22 (m, 2H), 1.76 (s, 3H), 2.13 (s, 3H), 2.12- 2.28 (m, 2H), 2.35- 2.41 (m, 2H), 3.37 (d, 2H, $J=7$ Hz), 3.71 (s, 3H), 4.22- 4.28 (m, 2H), 5.07 (s, 2H), 5.13- 5.17 (m, 1H) ppm; MS (m/z) 419.3 $[\text{M}-\text{H}]^-$, 443.2 $[\text{M}+\text{Na}]^+$.



((6-[6-Methoxy-7-methyl-3-oxo-4-(2-trimethylsilanyl-ethoxy)-1,3-dihydro-isobenzofuran-5-yl]-4-methyl-hex-4-enoylamino)-methyl)-phosphonic acid diethyl ester

To a solution of 6-[6-methoxy-7-methyl-3-oxo-4-(2-trimethylsilanyl-ethoxy)-1,3-dihydro-isobenzofuran-5-yl]-4-methyl-hex-4-enoic acid (50 mg, 0.12 mmol) in THF (1 mL) was added isobutyl chloroformate (17 μL , 0.13 mmol) and triethylamine (50 μL , 0.36 mmol) at 0°C . After stirring at 0°C for 2 hours, diethyl (aminomethyl) phosphonate oxalate (62 mg, 0.26 mmol) was added and stirring was continued at room temperature for 20 minutes. After removal of solvent, the residue was purified by preparative reverse-phase HPLC to afford 54.8 mg (81%) of the desired product. ^1H NMR (300 MHz, CDCl_3) δ 0.03 (s, 9H), 1.15- 1.22 (m, 2H), 1.31 (t, 6H), 1.81 (s, 3H), 2.18 (s, 3H), 2.30 (m, 4H), 3.41 (d, 2H, $J=7$ Hz), 3.65 (dd, 2H, $J=6, 12$ Hz), 3.77 (s, 3H), 3.77-4.16 (m, 4H), 4.26-4.32 (m, 2H), 5.12 (s, 2H), 5.17- 5.19 (m, 1H), 5.86 (bs, 1H) ppm; ^{31}P (121.4 MHz, CDCl_3) δ 23.01 ppm; MS (m/z) 568 $[\text{M}-\text{H}]^-$, 592 $[\text{M}+\text{Na}]^+$.

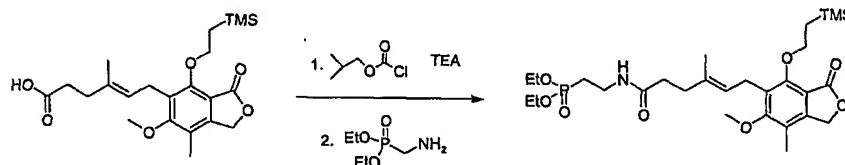


{[6-(4-Hydroxy-6-methoxy-7-methyl-3-oxo-1,3-dihydro-isobenzofuran-5-yl)-4-methyl-hex-4-enoylamino]-methyl}-phosphonic acid

To a solution of ({6-[6-methoxy-7-methyl-3-oxo-4-(2-trimethylsilyl-ethoxy)-1,3-dihydro-isobenzofuran-5-yl]-4-methyl-hex-4-enoylamino}-methyl)-phosphonic acid diethyl ester (40 mg, 0.07 mmol) in acetonitrile (1 mL) was added TMSBr (91 μ L, 0.7 mmol) followed by 2,6-lutidine (81.5 μ L, 0.7 mmol). The reaction was allowed to proceed overnight when it was completed as judged by LCMS. The reaction mixture was quenched with MeOH and concentrated to dryness. The residue was purified by preparative reverse-phase HPLC to afford 2.6 mg (9%) of desired product as a white solid. ^1H NMR (300 MHz, CD_3OD) δ 1.67 (s, 3H), 2.17 (m, 5H), 2.30-2.46 (m, 2H), 2.80- 2.86 (m, 2H), 3.55 (m, 2H), 3.82 (s, 3H), 5.26 (s, 3H) ppm; ^{31}P (121.4 MHz, CD_3OD) δ 10.27 ppm; MS (m/z) 412 $[\text{M}-\text{H}]^-$, 414 $[\text{M}+\text{H}]^+$.

Example 322

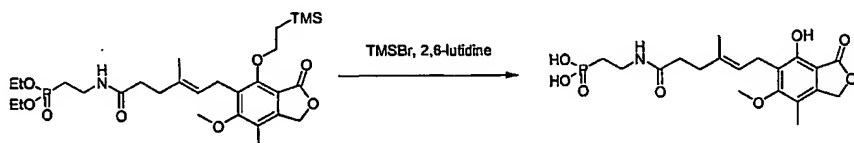
Representative compounds of the invention can be prepared as illustrated below.



(2-{6-[6-Methoxy-7-methyl-3-oxo-4-(2-trimethylsilyl-ethoxy)-1,3-dihydro-isobenzofuran-5-yl]-4-methyl-hex-4-enoylamino}-ethyl)-phosphonic acid diethyl ester

To a solution of 6-[6-methoxy-7-methyl-3-oxo-4-(2-trimethylsilyl-ethoxy)-1,3-dihydro-isobenzofuran-5-yl]-4-methyl-hex-4-enoic acid (50 mg, 0.12 mmol) in THF (1 mL) was added isobutyl chloroformate (17 μ L, 0.13 mmol) and triethylamine (50 μ L, 0.36 mmol) at 0°C. After stirring at 0°C for 2 hours, diethyl (aminoethyl) phosphonate oxalate (62 mg, 0.26 mmol) was added and stirred at room temperature was continued for one hour. After removal of solvent, the residue was purified by preparative reverse-phase HPLC to afford 37

mg (54%) of the desired product as a white solid. ^1H NMR (300 MHz, CDCl_3) δ 0.03 (s, 9H), 1.15-1.22 (m, 2H), 1.31 (t, 6H), 1.81 (s, 3H), 1.85-1.93 (m, 2H), 2.18 (s, 3H), 2.30 (m, 4H), 3.41 (d, 2H, $J=7$ Hz), 3.48-3.54 (m, 2H), 3.77 (s, 3H), 3.77-4.16 (m, 4H), 4.26-4.32 (m, 2H), 5.12 (s, 2H), 5.17-5.19 (m, 1H), 6.30 (bs, 1H) ppm; ^{31}P (121.4 MHz, CDCl_3) δ 29.91 ppm; MS (m/z) 584 $[\text{M}+\text{H}]^+$.

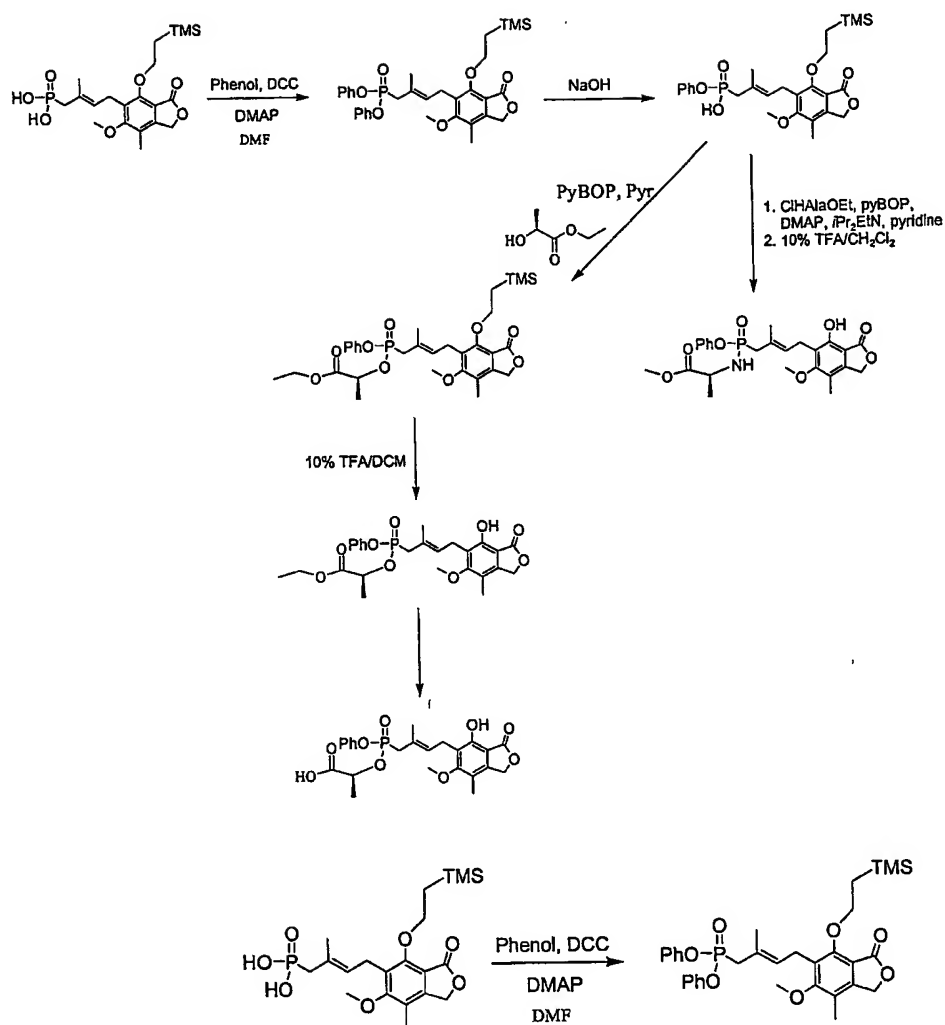


{2-[6-(4-Hydroxy-6-methoxy-7-methyl-3-oxo-1,3-dihydro-isobenzofuran-5-yl)-4-methyl-hex-4-enoylamino]-ethyl}-phosphonic acid

To a solution of (2-{6-[6-methoxy-7-methyl-3-oxo-4-(2-trimethylsilyl-ethoxy)-1,3-dihydro-isobenzofuran-5-yl]-4-methyl-hex-4-enoylamino}-ethyl)-phosphonic acid diethyl ester (36.6 mg, 0.063 mmol) in acetonitrile (1 mL) was added TMSBr (81 μL , 0.63 mmol) followed by 2,6-lutidine (73 μL , 0.63 mmol). The reaction was allowed to proceed overnight, when it was completed as judged by LCMS. The reaction mixture was quenched with MeOH and concentrated to dryness. The residue was purified by preparative reverse-phase HPLC to afford 5.8 mg (29%) of desired product as a white solid. ^1H NMR (300 MHz, CD_3OD) δ 1.80 (s, 3H), 2.14 (m, 5H), 2.25 (m, 4H), 3.35 (m, 2H), 3.38-3.38 (m, 2H), 3.75 (s, 3H), 5.23 (s, 3H) ppm; ^{31}P (121.4 MHz, CD_3OD) δ 26.03 ppm; MS (m/z) 426 $[\text{M}-\text{H}]^-$, 428 $[\text{M}+\text{H}]^+$.

Example 323

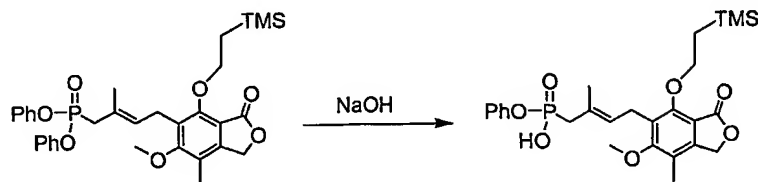
Representative compounds of the invention can be prepared as illustrated below.



{4-[6-Methoxy-7-methyl-3-oxo-4-(2-trimethylsilylanyl-ethoxy)-1,3-dihydro-isobenzofuran-5-yl]-2-methyl-but-2-enyl}-phosphonic acid diphenyl ester

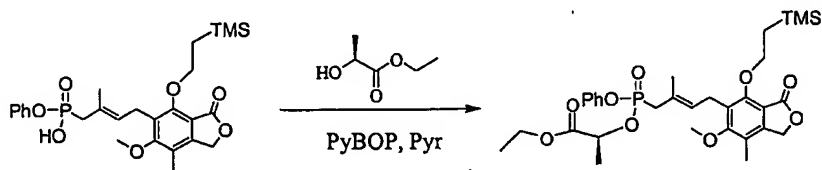
To a solution of [4-[6-methoxy-7-methyl-3-oxo-4-(2-trimethylsilylanyl-ethoxy)-1,3-dihydro-isobenzofuran-5-yl]-2-methyl-but-2-enyl}-phosphonic acid (260 mg, 0.59 mmol) in DMF (6 mL) and phenol (555 mg, 5.9 mmol) was added dicyclohexyl carbodiimide (1.21 g, 5.9 mmol) and DMAP (36 mg, 0.295 mmol). The reaction mixture was heated to 140°C for 30 minutes. After cooling to room temperature, the mixture was partitioned between EtOAc/Hexane (1:1) and 5% aqueous LiCl solution. The organic layer was washed with 5% aqueous LiCl solution repeatedly, then dried over Na_2SO_4 . After removal of solvent, the

residue was purified by silica gel chromatography to provide 75 mg (21%) of the desired product. MS (m/z) 617 $[M+Na]^+$.



{4-[6-Methoxy-7-methyl-3-oxo-4-(2-trimethylsilyl-ethoxy)-1,3-dihydro-isobenzofuran-5-yl]-2-methyl-but-2-enyl}-phosphonic acid monophenyl ester

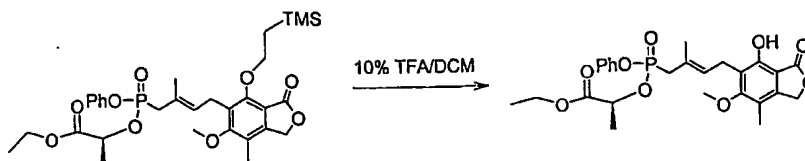
To a solution of {4-[6-methoxy-7-methyl-3-oxo-4-(2-trimethylsilyl-ethoxy)-1,3-dihydro-isobenzofuran-5-yl]-2-methyl-but-2-enyl}-phosphonic acid diphenyl ester (75 mg, 0.126 mmol) in THF (5 mL) was added 1N NaOH (0.1 mL) solution. The mixture was allowed to stir at room temperature for 16 hours. EtOAc was added and the resulting mixture was washed with 1N HCl. The organic layer was concentrated to dryness and the residue was purified by RP HPLC using a C18 column with a gradient of H₂O, 0.1% TFA-acetonitrile, 0.1% TFA to provide 24.8 mg (38 %) of the desired product. MS (m/z) 517 $[M-H]^-$, 541 $[M+Na]^+$.



2-({4-[6-Methoxy-7-methyl-3-oxo-4-(2-trimethylsilyl-ethoxy)-1,3-dihydro-isobenzofuran-5-yl]-2-methyl-but-2-enyl}-phenoxy-phosphinoyloxy)-propionic acid ethyl ester

To a solution of {4-[6-methoxy-7-methyl-3-oxo-4-(2-trimethylsilyl-ethoxy)-1,3-dihydro-isobenzofuran-5-yl]-2-methyl-but-2-enyl}-phosphonic acid monophenyl ester (25 mg, 0.048 mmol) and ethyl (S)-(-)-lactate (34 mg, 0.288 mmol) in pyridine (1 mL) was added PyBOP (125 mg, 0.24 mmol). The solution was stirred at room temperature for 16 hours and concentrated. The

residue was purified by RP HPLC using a C18 column with a gradient of H₂O, 0.1% TFA-acetonitrile, 0.1% TFA to provide 24 mg (83%) of the desired product. MS (*m/z*) 641 [M+Na]⁺.

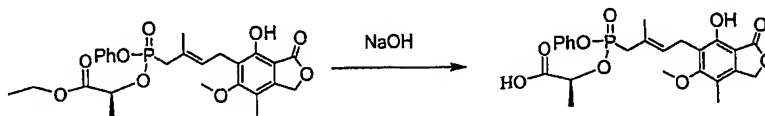


2-{{[4-(4-Hydroxy-6-methoxy-7-methyl-3-oxo-1,3-dihydro-isobenzofuran-5-yl)-2-methyl-but-2-enyl]-phenoxy-phosphinoyloxy}-propionic acid ethyl ester

To a solution of 2-({[4-[6-methoxy-7-methyl-3-oxo-4-(2-trimethylsilyl-ethoxy)-1,3-dihydro-isobenzofuran-5-yl]-2-methyl-but-2-enyl]-phenoxy-phosphinoyloxy)-propionic acid ethyl ester (24 mg, 0.039 mmol) in DCM (1 mL) was added TFA (0.5 mL) and the mixture was stirred at room temperature for 10 minutes. The reaction mixture was dried under reduced pressure and the residue was purified by RP-HPLC to provide 18 mg (90%) of the desired product as a clear oil. ¹H NMR (300 MHz, CDCl₃) δ 1.18-1.34 (m, 3H), 1.36-1.48 (dd, 3H), 2.02 (m, 3H), 2.17 (s, 3H), 2.78-2.98 (dd, 2H), 3.45 (m, 2H), 3.79 (s, 3H), 4.05-4.25 (m, 2H), 4.97 (m, 1H), 5.21 (s, 2H), 5.48 (t, *J* = 7.2 Hz, 1H), 7.05-7.18 (m, 5H) ppm; ³¹P (121.4 MHz, CDCl₃) δ 24.59, 26.13 ppm; MS (*m/z*) 517 [M-H]⁻, 519 [M+H]⁺.

Example 324

Representative compounds of the invention can be prepared as illustrated below.

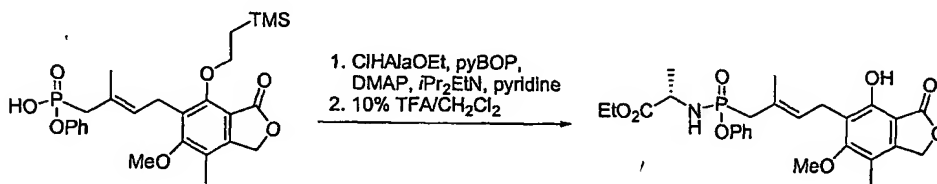


2-{{[4-(4-Hydroxy-6-methoxy-7-methyl-3-oxo-1,3-dihydro-isobenzofuran-5-yl)-2-methyl-but-2-enyl]-phenoxy-phosphinoyloxy}-propionic acid

To a solution of 2-[[4-(4-hydroxy-6-methoxy-7-methyl-3-oxo-1,3-dihydro-isobenzofuran-5-yl)-2-methyl-but-2-enyl]-phenoxy-phosphinoyloxy]-propionic acid ethyl ester (10 mg, 0.019 mmol) in THF (3 mL) was added 1N NaOH (232 μ L), and the mixture was stirred at room temperature for 1 hour. The reaction mixture was dried under reduced pressure and the residue was purified by RP-HPLC to provide 6 mg (77 %) of the desired product as a clear oil. ^1H NMR (300 MHz, CD_3OD) δ 1.41 (d, $J = 7$ Hz, 3H), 1.97 (s, 3H), 2.16 (s, 3H), 2.59 (d, $J = 22$ Hz, 2H), 3.45 (m, 2H), 3.79 (s, 3H), 4.83 (m, 1H), 5.26 (s, 2H), 5.43 (t, $J = 7.2$ Hz, 1H) ppm; ^{31}P (121.4 MHz, CD_3OD) δ 27.02 ppm; MS (m/z) 413 $[\text{M}-\text{H}]^-$, 415 $[\text{M}+\text{H}]^+$.

Example 325

Representative compounds of the invention can be prepared as illustrated below.



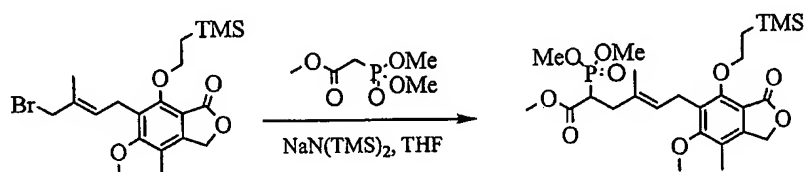
2-[[4-(4-Hydroxy-6-methoxy-7-methyl-3-oxo-1,3-dihydro-isobenzofuran-5-yl)-2-methyl-but-2-enyl]-phenoxy-phosphinoylamino]-propionic acid ethyl ester

{4-[6-methoxy-7-methyl-3-oxo-4-(2-trimethylsilyl-ethoxy)-1,3-dihydro-isobenzofuran-5-yl]-2-methyl-but-2-enyl}-phosphonic acid monophenyl ester (1 g, ~ 1.9 mmol) was combined with pyBOP (2 g, 4 mmol) and DMAP (120 mg, 0.96 mmol). A solution of L-alanine ethyl ester hydrochloride salt (2.9 g, 19 mmol) and diisopropylethylamine (6.7 mL, 38 mmol) in pyridine (5 mL) was added to the monoacid mixture and the reaction was stirred at room temperature for 12 hours. The reaction mixture was then concentrated and purified twice by column chromatography (1% MeOH/ CH_2Cl_2 3% MeOH/ CH_2Cl_2). The resulting oil was dissolved in a vigorously-stirred solution of 10% TFA/ CH_2Cl_2 (30 mL) at -40°C . The reaction was gradually warmed to

0°C. After about 3 hours, the reaction was complete. Pyridine (4.5 mL) was added, and the reaction mixture was concentrated. The product was purified by preparative TLC (5% MeOH/CH₂Cl₂) and concentrated to give 210 mg (21%) of the desired product as a light yellow oil. ¹H NMR (300 MHz, CDCl₃) δ 7.83-7.70 (m, 1H), 7.30-7.20 (m, 2H), 7.18-7.03 (m, 3H), 5.60-5.35 (m, 1H), 5.21 (s, 2H), 4.17-3.95 (m, 3H), 3.79 (s, 3H), 3.60-3.40 (m, 3H), 2.80-2.60 (m, 2H), 2.17 (m, 3H), 2.01 (m, 3H), 1.30-1.10 (m, 6H) ppm; ³¹P NMR (121 MHz, CDCl₃) δ 28.0, 27.5 ppm; MS (*m/z*) 516 [M-H]⁻.

Example 326

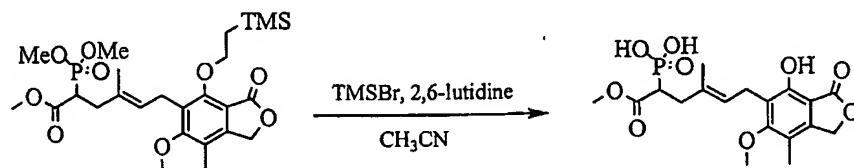
Representative compounds of the invention can be prepared as illustrated below.



2-(Dimethoxy-phosphoryl)-6-[6-methoxy-7-methyl-3-oxo-4-(2-trimethylsilyl-ethoxy)-1,3-dihydro-isobenzofuran-5-yl]-4-methyl-hex-4-enoic acid methyl ester

To a solution of trimethylphosphonoacetate (63 μL, 0.39 mmol) in THF (1 mL) was added NaN(TMS)₂ (0.39 mmol, 0.39 mL) at ambient temperature. After 30 minutes, a solution of 6-(4-bromo-3-methyl-but-2-enyl)-5-methoxy-4-methyl-7-(2-trimethylsilyl-ethoxy)-3H-isobenzofuran-1-one (69 mg, 0.156 mmol) in THF (1 mL) was added. The reaction mixture was stirred for 2 hours, when a precipitate was observed. The reaction mixture was worked up by addition of a saturated aqueous solution of ammonium chloride and extraction of the product with EtOAc. The organic extract was dried and the product was purified using silica gel chromatography with 0-100% EtOAc-Hexanes to provide 40 mg of the desired product as a colorless oil. ¹H NMR (300 MHz, CDCl₃) δ 0.05 (s, 9H), 1.20-1.26 (m, 2H), 1.79 (s, 3H), 2.17 (s, 3H), 2.42-2.72 (m, 2H), 3.19 (ddd, 1H, *J* = 4, 12, 23 Hz), 3.39 (d, 2H, *J* = 7 Hz), 3.62 (s, 3H),

3.75 (s, 3H), 3.77- 3.84 (m, 6H), 4.27- 4.34 (m, 2H), 5.12 (s, 2H), 5.24 (t, 1H, $J=7$ Hz) ppm; ^{31}P (121.4 MHz, CDCl_3) δ 25.1 ppm; MS (m/z) 565.2 $[\text{M}+\text{Na}]^+$.

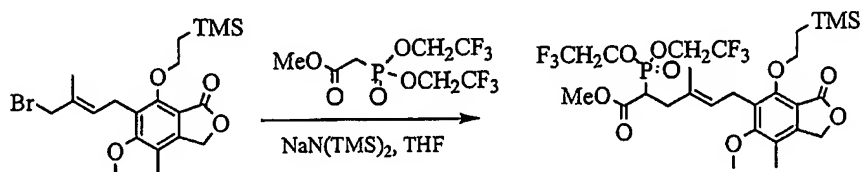


6-(4-Hydroxy-6-methoxy-7-methyl-3-oxo-1,3-dihydro-isobenzofuran-5-yl)-4-methyl-2-phosphono-hex-4-enoic acid methyl ester

To a solution of 2-(dimethoxy-phosphoryl)-6-[6-methoxy-7-methyl-3-oxo-4-(2-trimethylsilyl-ethoxy)-1,3-dihydro-isobenzofuran-5-yl]-4-methyl-hex-4-enoic acid methyl ester (30 mg, 0.055 mmol) in acetonitrile (2 mL) was added trimethylsilyl bromide (0.18 mL). After 10 minutes, 2,6-lutidine (0.16 mL) was added to the reaction at ambient temperature. The reaction was allowed to proceed for 16 hours before it was concentrated to dryness. The residue was resuspended in a solution of DMF: H_2O (8: 2, 1 mL) and purified by RP HPLC using a C18 column with a gradient of H_2O , 0.1% TFA-acetonitrile, 0.1% TFA to provide 18 mg of the product as a white powder. ^1H NMR (300 MHz, CD_3OD) δ 1.81 (s, 3H), 2.16 (s, 3H), 2.40-2.49 (m, 1H), 2.63 (dt, 1H, $J=6, 17$ Hz), 3.07 (ddd, 1H, $J=4, 12, 23$ Hz), 3.38 (3, 2H, $J=7$ Hz), 3.52 (s, 3H), 3.77 (s, 3H), 5.25 (s, 2H), 5.28 (t, 1H, $J=7$ Hz) ppm; ^{31}P (121.4 MHz, CDCl_3) δ 19.5 ppm; MS (m/z) 415.2 $[\text{M}+\text{H}]^+$, 437.2 $[\text{M}+\text{Na}]^+$.

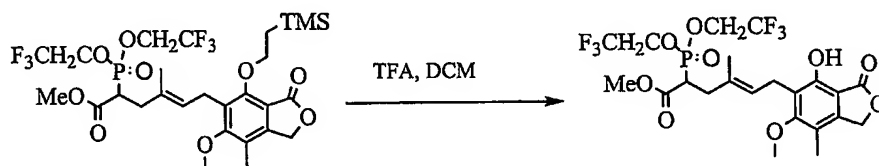
Example 327

Representative compounds of the invention can be prepared as illustrated below.



2-(Bis-(2,2,2-trifluoroethoxy)phosphoryl)-6-[6-methoxy-7-methyl-3-oxo-4-(2-trimethylsilanyl-ethoxy)-1,3-dihydro-isobenzofuran-5-yl]-4-methyl-hex-4-enoic acid methyl ester

To a solution of [bis-(2,2,2-trifluoro-ethoxy)-phosphoryl]-acetic acid methyl ester (186 μ L, 0.88 mmol) in anhydrous THF (2 mL) was added a solution of 1N NaN(TMS)₂ in THF (0.88 mL, 0.88 mmol). The solution was stirred at room temperature for 30 minutes, whereupon a solution of 6-(4-bromo-3-methyl-but-2-enyl)-5-methoxy-4-methyl-7-(2-trimethylsilanyl-ethoxy)-3H-isobenzofuran-1-one (98 mg, 0.22 mmol) in THF (1 mL) was added. The reaction mixture was stirred overnight when a precipitate was observed. The reaction mixture was worked up by addition of a saturated aqueous solution of ammonium chloride and extraction of the product with EtOAc. The organic extract was dried and the product was purified by RP HPLC using a C18 column with a gradient of H₂O, 0.1% TFA-acetonitrile, 0.1% TFA to provide 72 mg (48%) of the product as a colorless oil. ¹H NMR (300 MHz, CDCl₃) δ 0.05 (s, 9H), 1.22 (t, 3H, *J* = 7 Hz), 1.81 (s, 3H), 2.18 (s, 3H), 2.5- 2.7 (m, 2H), 3.3 (ddd, 1H, *J* = 4, 12, 23 Hz), 3.40 (d, 2H, *J* = 7 Hz), 3.65 (s, 3H), 3.76 (s, 3H), 4.29- 5.13 (m, 6H), 5.13 (s, 2H), 5.28 (t, 1H, *J* = 7 Hz) ppm; MS (*m/z*) 701.2 [M+Na]⁺.



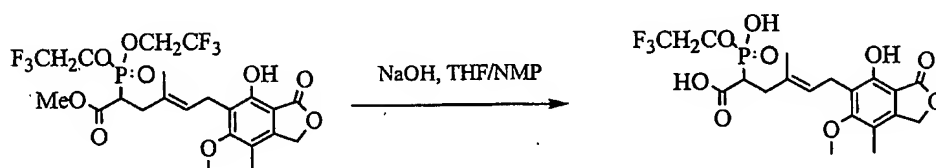
2-(Bis-(2,2,2-trifluoroethoxy)phosphoryl)-6-[6-methoxy-7-methyl-3-oxo-4-(2-hydroxyoxy)-1,3-dihydro-isobenzofuran-5-yl]-4-methyl-hex-4-enoic acid methyl ester

[2-(Bis-(2,2,2-trifluoroethoxy)phosphoryl)-6-[6-methoxy-7-methyl-3-oxo-4-(2-trimethylsilanyl-ethoxy)-1,3-dihydro-isobenzofuran-5-yl]-4-methyl-hex-4-enoic acid methyl ester (70 mg) was dissolved in a solution of 10% trifluoroacetic acid in dichloromethane (5mL). After 10 minutes, the mixture was concentrated and the product was purified by RP HPLC using a C18 column with a gradient of H₂O, 0.1% TFA-acetonitrile, 0.1% TFA to provide 45 mg

(75%) of the product as a colorless oil. ^1H NMR (300 MHz, CDCl_3) δ 1.81 (s, 3H), 2.16 (s, 3H), 2.5- 2.7 (m, 2H), 3.3 (ddd, 1H), 3.38 (d, 2H, $J = 7$ Hz), 3.65 (s, 3H), 3.77 (s, 3H), 4.33- 4.43 (m, 4H), 5.21 (s, 2H), 5.33 (t, 1H, $J = 7$ Hz) ppm; ^{31}P (121.4 MHz, CDCl_3) δ 25.8 ppm; MS (m/z) 601.2 $[\text{M}+\text{Na}]^+$.

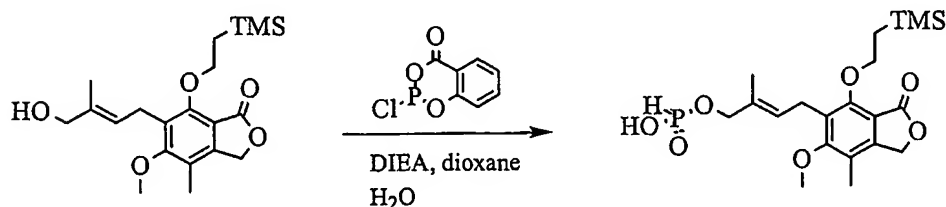
Example 328

Representative compounds of the invention can be prepared as illustrated below.



6-(4-Hydroxy-6-methoxy-7-methyl-3-oxo-1,3-dihydro-isobenzofuran-5-yl)-2-[hydroxy-(2,2,2-trifluoro-ethoxy)-phosphoryl]-4-methyl-hex-4-enoic acid

To a solution of [bis-(2,2,2-trifluoro-ethoxy)-phosphoryl]-acetic acid methyl ester (186 μL , 0.88 mmol) in anhydrous THF (0.5 mL) was added a solution of 1N NaOH (aqueous; 0.06 mL) and N-methylpyrrolidinone (0.2mL). After 6.5 hours, another aliquot of 1N NaOH (0.06mL) was added and the mixture was stirred overnight. After concentration, the residue was suspended in DMF (<1mL), neutralized with a few drops of TFA and purified by RP HPLC using a C18 column with a gradient of H_2O , 0.1% TFA-acetonitrile, 0.1% TFA to provide 5.6 mg (72%) of the product as a white powder after lyophilization. ^1H NMR (300 MHz, CD_3OD) δ 1.83 (s, 3H), 2.16 (s, 3H), 2.43-2.51 (m, 1H), 2.59-2.70 (m, 1H), 3.13 (ddd, 1H), 3.40 (d, 2H), 3.76 (s, 3H), 4.36-4.47 (m, 2H), 5.25 (s, 2H), 5.34 (t, 1H, $J = 7$ Hz) ppm; MS (m/z) 505.2 $[\text{M}+\text{Na}]^+$.

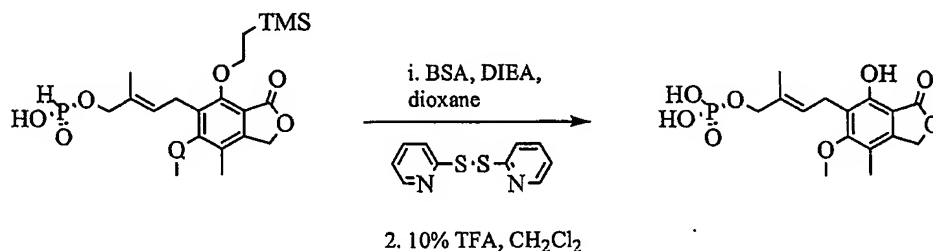


Example 329

Representative compounds of the invention can be prepared as illustrated below.

Phosphorous acid mono-{4-[6-methoxy-7-methyl-3-oxo-4-(2-trimethylsilyl-ethoxy)-1,3-dihydro-isobenzofuran-5-yl]-2-methyl-but-2-enyl} ester

To a solution of 6-(4-hydroxy-3-methyl-but-2-enyl)-5-methoxy-4-methyl-7-(2-trimethylsilyl-ethoxy)-3H-isobenzofuran-1-one (75 mg, 0.20 mmol) and DIEA (49 μ L, 0.28 mmol) in dioxane (2 mL) was added 2-chloro-4H-1,3,2-benzodioxaphosphorin-4-one (56.7 mg, 0.28 mmol) according the procedure of Shadid, B. et al., *Tetrahedron*, **1989**, 45, 12, 3889. After 10 minutes, another portion of 2-chloro-4H-1,3,2-benzodioxaphosphorin-4-one (40 mg, 0.20 mmol) and DIEA (35 μ L, 0.20 mmol) were added. The reaction was allowed to proceed at room temperature for an additional hour, after which it was quenched by the addition of H₂O. The solution was stirred for another 10 minutes and concentrated *in vacuo* to a small volume. The product was triturated with diethyl ether and coevaporated from acetonitrile (4 x 10 mL) to provide the product. ¹H NMR (300 MHz, CDCl₃) δ 0.03 (s, 9H), 1.08- 1.30 (m, 2H), 1.84 (br s, 3H), 2.17 (s, 3H), 3.46 (br s, 2H), 3.76 (s, 3H), 4.21- 4.39 (m, 4H), 5.12 (s, 2H), 5.43- 5.60 (m, 1H), 7.83 (br s, 1H); ³¹P (121.4 MHz, CDCl₃) δ 7.22; MS (*m/z*) 441 [M-H].

**Example 330**

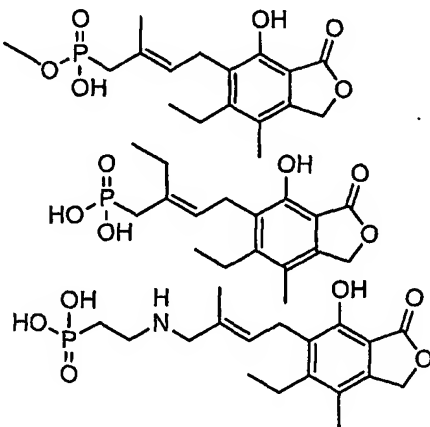
Representative compounds of the invention can be prepared as illustrated below.

Phosphoric acid mono-{4-[6-methoxy-7-methyl-3-oxo-4-(2-trimethylsilylanyl-ethoxy)-1,3-dihydro-isobenzofuran-5-yl]-2-methyl-but-2-enyl} ester

A solution of phosphorous acid mono-{4-[6-methoxy-7-methyl-3-oxo-4-(2-trimethylsilylanyl-ethoxy)-1,3-dihydro-isobenzofuran-5-yl]-2-methyl-but-2-enyl} ester (27 mg, 0.06 mmol) in dioxane (1 mL) was stirred with DIEA (21 μ L, 0.12 mmol) and N,O-bis(trimethylsilyl)acetamide (29 μ L, 0.12 mmol) at room temperature for 3 hours. To the reaction solution was added 2,2'-dipyridyldisulfide (16 mg, 0.072 mmol) and the mixture was allowed to stir for an additional 2 hours at room temperature. The reaction mixture was diluted by addition of H₂O and the solution was stirred for 2 more hours when it was concentrated. The residue was dissolved in a solution of 10% TFA/ CH₂Cl₂ and stirred at room temperature for 9 hours. The reaction mixture was dried under reduced pressure and the product was purified by reverse-phase HPLC to provide the desired product as a white solid. ¹H NMR (300 MHz, CD₃OD) δ 1.87 (s, 3H), 2.16 (s, 3H), 3.47 (d, 2H, *J*= 7 Hz), 3.79 (s, 3H), 4.28 (d, 2H, *J*= 6 Hz), 5.26 (s, 2H), 5.50- 5.61 (m, 1H); ³¹P (121.4 MHz, CD₃OD) δ 0.50; MS (*m/z*) 357 [M-H]⁻.

Example 331

Several compounds of the invention are presented below.



Example 332

Additional representative compounds of the invention, and intermediates thereof, can be prepared according to the methods presented below.



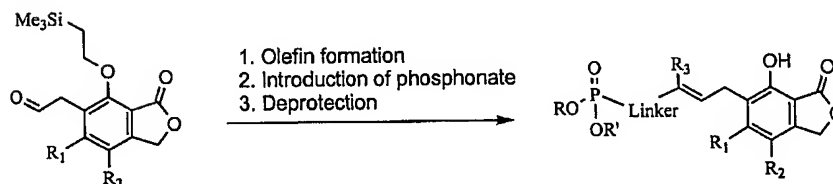
Linker = 0-8 atoms, preferably 1-6;

R_1 = OMe, OEt, vinyl, Et, cyclopropyl, NHMe, NHCHO

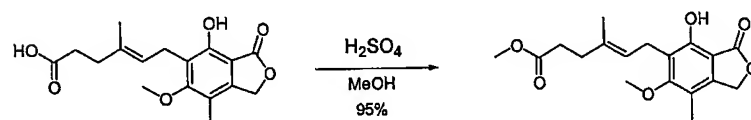
R_2 = Me, Cl, CF_3

R_3 = H, Me, cyclopropyl, Et, vinyl, CF_3

R_4 = H, Cl, Me, Et, cyclopropyl, vinyl, allyl,

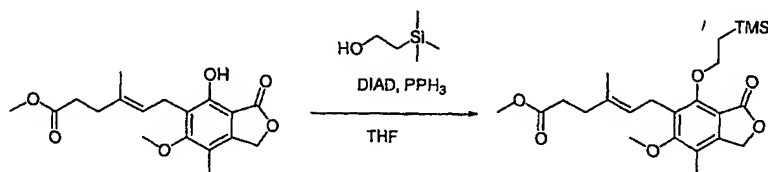
**Synthesis of phenacetaldehydes with variants at R_1 , R_2**

The parent compound (R_1 = OMe; R_2 = Me) is accessible by semi-synthesis from mycophenolic acid as follows:

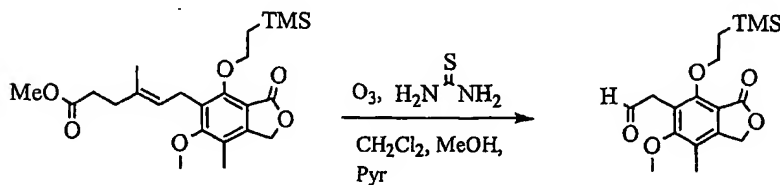


To a solution of mycophenolic acid (500 g, 1.56 mol) in MeOH (4 L) under nitrogen atmosphere was added sulfuric acid (10 mL) dropwise, and the suspension was stirred at room temperature. After 2 hours, the reaction became homogeneous, and soon thereafter a precipitate was formed. The reaction was allowed to stir at room temperature for 10 hours, at which time TLC indicated complete reaction. The reaction was cooled in an ice bath to 10°C and then

filtered using a Buchner funnel. The filter cake was washed with ice cold methanol (750 mL) followed by hexanes (750 mL) and then dried to give 497 g (95%) of the desired product as a solid: ^1H NMR (300 MHz, CDCl_3) δ , 1.81 (s, 3H), 2.18 (s, 3H), 2.15 (s, 3H), 2.37-2.50 (m, 4H), 3.38 (d, 2H, $J=7$ Hz), 3.62 (s, 3H), 3.77 (s, 3H), 5.13 (s, 2H), 5.22 (m, 1H), 7.17 (s, 1H).

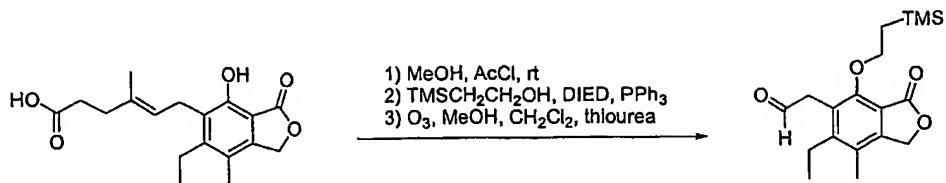


To a solution (3.99 g, 11.9 mmol), PPh_3 (4.68 g, 17.9 mmol), and diisopropyl azodicarboxylate (3.46 mL, 17.9 mmol) in THF (60 mL) at 0°C was added a solution of 2-trimethylsilylethanol (2.05 mL, 14.3 mmol) in THF (20 mL). The resulting yellow solution was allowed to warm to room temperature and stirred for 4 hours. The reaction was worked up by concentrating the solution to dryness and addition of ether and hexanes. Triphenylphosphine oxide was removed by filtration and the filtrate was concentrated and purified by silica gel chromatography to provide 4.8 g (100%) as a clear oil: ^1H NMR (300 MHz, CDCl_3) δ 0.03 (s, 9H), 1.18-1.30 (m, 2H), 1.81 (s, 3H), 2.18 (s, 3H), 2.25-2.33 (m, 2H), 2.37-2.45 (m, 2H), 3.42 (d, 2H, $J=7$ Hz), 3.62 (s, 3H), 3.77 (s, 3H), 4.25-4.35 (m, 2H), 5.13 (s, 2H), 5.12-5.22 (m, 1H).

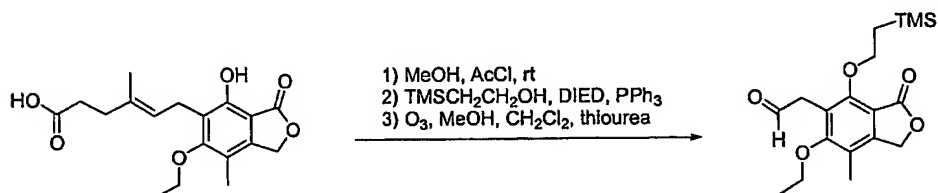


A solution (9.6 g, 22 mmol) in MeOH (90 mL), CH_2Cl_2 (90 mL) and pyridine (0.7 mL) was cooled to -70°C using a dry ice/acetone bath. A stream of ozone was bubbled through the reaction *via* a gas dispersion tube until the reaction became blue in color (1.5 hours). The ozone line was replaced with a stream of nitrogen and bubbling continued for another 30 minutes, by which

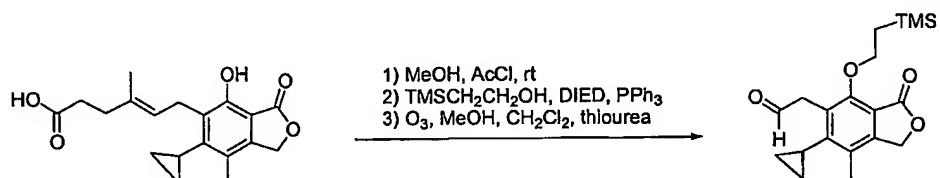
time the blue color had disappeared. To this solution at -70°C was added thiourea (1.2 g, 15.4 mmol) in one portion, and the cooling bath was removed. The reaction was allowed to warm to room temperature and stirred for 15 hours. The reaction was worked up by filtration to remove solid thiourea S-dioxide, and then partitioned between CH_2Cl_2 and water. The organic layer was removed. The aqueous layer was washed with CH_2Cl_2 and the organic extracts were combined, washed with aqueous 1N HCl, saturated NaHCO_3 and brine, and dried *in vacuo*. The residue was purified by silica gel chromatography to afford 7.3 g (99 %) as a white solid: ^1H NMR (300 MHz, CDCl_3) δ -0.01 (s, 9H), 1.05-1.15 (m, 2H), 2.15 (s, 3H), 3.69 (s, 3H), 3.78 (d, 2H, $J=1$ Hz), 4.27-4.39 (m, 2H), 5.11 (s, 2H), 9.72 (d, 1H, $J=1$ Hz).

R₁ variants

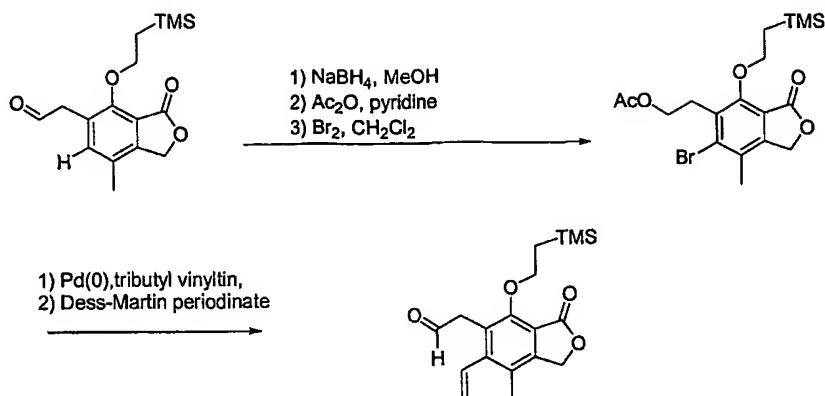
The starting material, synthesized according to *J. Med. Chem.*, 1996, 39, 4181-4196, is transformed to the desired aldehyde using methods analogous to those described above.



The starting material, synthesized according to *J. Med. Chem.*, 1996, 39, 4181-4196, is transformed to the desired aldehyde using methods analogous to those described above.



The starting material, synthesized according to *J. Med. Chem.*, 1996, 39, 4181-4196, is transformed to the desired aldehyde using methods analogous to those described above.



The aldehyde is dissolved in an organic solvent such as methanol and sodium borohydride is added. At the end of the reaction, aqueous HCl solution is added and the solvent is removed *in vacuo*. Further purification is achieved by chromatography.

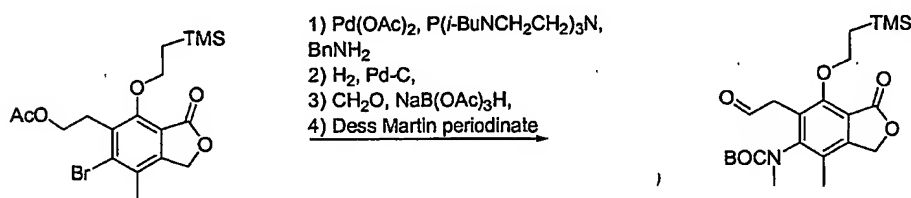
The resulting alcohol is dissolved in an organic solvent such as dichloromethane (DCM). Pyridine and acetic anhydride are added and stirring at room temperature is continued. At the end of the reaction additional DCM is added and the solution is washed with aqueous HCl solution, aqueous sodium bicarbonate solution, and dried over sodium sulfate. Filtration and evaporation of the solvent *in vacuo* gives the crude product. Further purification is achieved by chromatography.

The acetate is dissolved in DCM and bromine is added, according to a procedure from *J. Med. Chem.*, 1996, 39, 4181-4196. At the end of the reaction, additional DCM is added and the solution is washed with aqueous sodium thiosulfate solution and brine. The organic layer is dried over sodium sulfate. Filtration and evaporation of solvents yields the crude material. Further purification is achieved by chromatography.

The product of the previous step, lithium chloride, triphenylarsine, tributylvinyltin, and tris(dibenzylideneacetone)dipalladium(0)-chloroform adduct are heated in an organic solvent such as *N*-methylpyrrolidinone at an elevated temperature of approximately 55°C, according to a procedure from *J. Med. Chem.*, 1996, 39, 4181-4196. At the end of the reaction, the mixture is cooled to room temperature and poured into a mixture of ice, potassium fluoride, water, and ethyl acetate. Stirring is continued for one hour. The suspension is

filtered through Celite and extracted with ethyl acetate. The combined organic extracts are dried over sodium sulfate. The solvents are removed *in vacuo* and the crude material is further purified by chromatography.

The product of the previous step is dissolved in an organic solvent such as DCM or THF. 1,1,1-tris(aclyoxy)-1,1-dihydro-1,2benziodoxol-3-(1H)-one (Dess-Martin reagent) is added and the solution is stirred at room temperature, according to a procedure from *J. Org. Chem.*, 1984, 48, 4155-4156. At the end of the reaction diethyl ether is added, followed by aqueous sodium hydroxide solution. The layers are separated and the organic layer is washed with aqueous sodium hydroxide solution, water, and dried over sodium sulfate. Filtration and evaporation of solvents yields the crude product. Further purification is achieved by chromatography.



The starting material is dissolved in an organic solvent such as toluene. P(*isobutyl*NCH₂CH₂)₃N, palladium(II)acetate, sodium *tert.* butoxide, and benzylamine are added and the mixture was heated at 80°C, according to a procedure from *J. Org. Chem.*, 2003, 68, 452-459. At the end of the reaction, the mixture is cooled to room temperature and the solvents are removed *in vacuo*. The crude material is purified by chromatography. Any residual acetate is removed by brief treatment with methanolic sodium methoxide.

The benzyl-protected aniline is dissolved in an organic solvent such as DMF. Palladium on carbon is added and the reaction mixture is placed under an atmosphere of hydrogen. At the end of the reaction, the mixture is filtered through Celite. The solvents are removed *in vacuo*. Further purification is achieved by chromatography.

The resulting primary aniline is dissolved in an organic solvent such as THF, acetonitrile, or DMF and is treated with formaldehyde and sodium triacetoxyborohydride as described in *J. Org. Chem.*, 1996, 61, 3849-3862. The

reaction is quenched with aqueous sodium bicarbonate and the product is extracted with an organic solvent such as ethyl acetate. The crude material is treated with di-*t*-butyl dicarbonate in an organic solvent such as dimethylformamide and aqueous sodium hydroxide. The resulting carbamate is purified by chromatography.

The primary alcohol product is dissolved in an organic solvent such as DCM or THF. 1,1,1-tris(acyloxy)-1,1-dihydro-1,2-benziodoxol-3-(1H)-one (Dess-Martin reagent) is added and the solution is stirred at room temperature, according to a procedure from *J. Org. Chem.*, 1984, 48, 4155-4156. At the end of the reaction diethyl ether is added, followed by aqueous sodium hydroxide solution. The layers are separated and the organic layer is washed with aqueous sodium hydroxide solution, water, and dried over sodium sulfate. Filtration and evaporation of solvents yields the crude product. Further purification is achieved by chromatography.

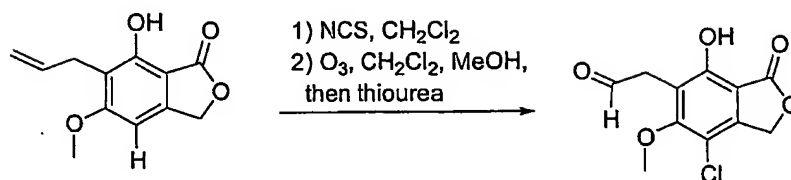


The starting material is dissolved in an organic solvent such as DCM or THF and is treated with the mixed anhydride of formic and pivalic acids, according to a procedure from *Recl. Trav. Chem. Pay-Bas*, 1982, 101, 460. At the end of the reaction, the solvent and all volatiles are removed *in vacuo* and the crude product is further purified by chromatography.

The product is dissolved in an organic solvent such as DCM or THF. 1,1,1-Tris(acyloxy)-1,1-dihydro-1,2-benziodoxol-3-(1H)-one (Dess-Martin reagent) is added and the solution was stirred at room temperature, according to a procedure from *J. Org. Chem.*, 1984, 48, 4155-4156. At the end of the reaction diethyl ether is added, followed by aqueous sodium hydroxide solution. The layers are separated and the organic layer is washed with aqueous sodium hydroxide solution, water, and dried over sodium sulfate. Filtration and

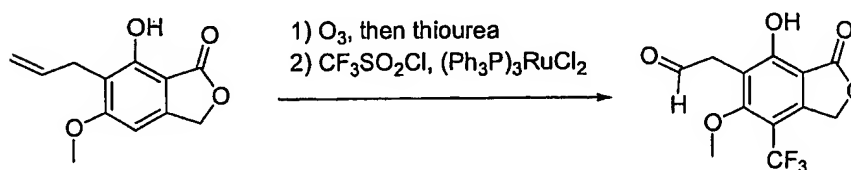
evaporation of solvents yields the crude product. Further purification is achieved by chromatography.

R₂ variants



The starting material is dissolved in an organic solvent such as DMF and reacted with *N*-chlorosuccinimide, according to a procedure from *J. Med. Chem.*, 1996, 39, 4181-4196. After the starting material is consumed the reaction mixture is poured into water and the product is extracted with diethyl ether. The combined organic layers are dried over sodium sulfate. Filtration and evaporation of the solvent yields a crude reaction product.

The product of step one is dissolved in a mixture of organic solvents such as methanol, DCM, and pyridine. The solution is cooled to -78°C and ozone is bubbled into the solution until a blue color persists. The excess ozone is removed with a nitrogen stream. The reaction mixture is warmed to room temperature and thiourea is added. Stirring at room temperature is continued. The reaction mixture is filtered and partitioned between DCM and water. The aqueous layer is extracted with DCM and the combined organic layers are washed with HCl (1 N), saturated aqueous sodium bicarbonate solution and brine. The solution is dried over sodium sulfate. Filtration and evaporation of the solvents yields the crude aldehyde. Further purification is achieved by chromatography.

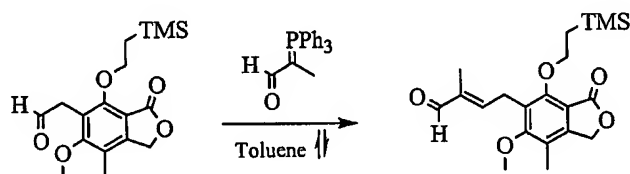


The starting material is dissolved in a mixture of organic solvents such as methanol, DCM, and pyridine. The solution is cooled to -78°C and ozone is

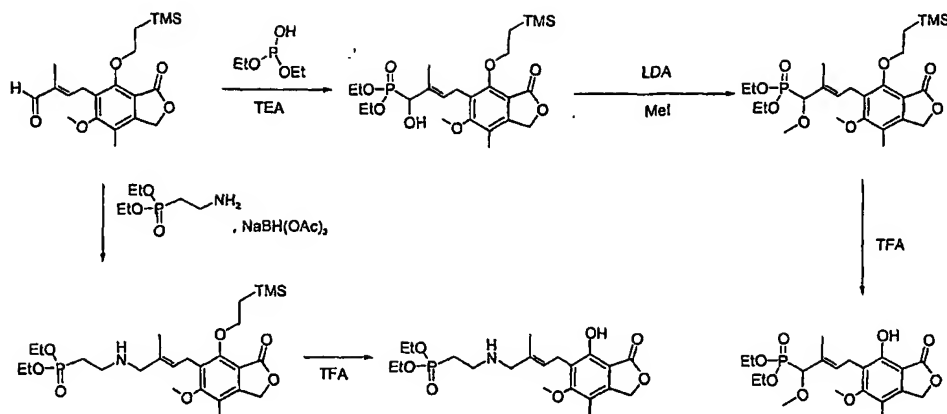
bubbled into the solution until a blue color persists. The excess ozone is removed with a nitrogen stream. The reaction mixture is warmed to room temperature and thiourea is added. Stirring at room temperature is continued. The reaction mixture is filtered and partitioned between DCM and water. The aqueous layer is extracted with DCM and the combined organic layers are washed with HCl (1 N), saturated aqueous sodium bicarbonate solution, and brine. The solution is dried over sodium sulfate. Filtration and evaporation of the solvents yields the crude aldehyde. Further purification is achieved by chromatography.

The product of step one is dissolved in an organic solvent such as benzene. Trifluoromethanesulfonyl chloride and dichlorotris(triphenylphosphine)rhuthenium are added and the solution is degassed. The reaction mixture is heated at 120 °C, according to a procedure from *J. Chem. Soc., Perkin Trans. 1*, 1994, 1339-1346. At the end of the reaction the mixture is cooled to room temperature and the solvent is removed *in vacuo*. Further product purification is achieved by chromatography.

Synthesis of olefins and linkers to phosphonates

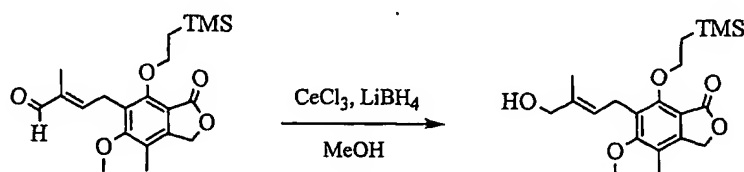


The phenacetaldehyde (5.3 g, 15.8 mmol) in toluene (50 mL) was heated at 100°C with 2-(triphenyl-phosphanylidene)-propionaldehyde (6.8 g, 20.5 mmol) overnight. After concentration, the residue was purified by silica gel chromatography to provide 4.24 g (72%) of the unsaturated aldehyde as a pale yellow solid. ¹H NMR (300 MHz, CDCl₃) δ 0.00 (s, 9H), 1.10- 1.21 (m, 2H), 1.87 (s, 3H), 2.16 (s, 3H), 3.67- 3.76 (m, 2H), 3.74 (s, 3H), 4.27- 4.39 (m, 2H), 5.11 (s, 2H), 6.40- 6.48 (m, 1H), 9.2 (s, 1H).



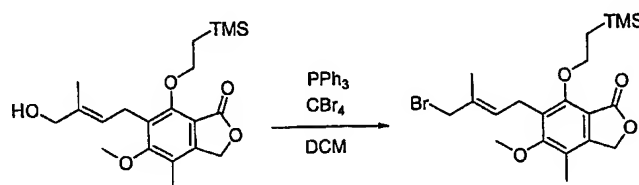
The trimethylsilyl-protected aldehyde is treated with diethylphosphite in a solvent such as acetonitrile in the presence of a base such as triethylamine to afford the hydroxy phosphonate, according to a procedure such as that reported in *Tetrahedron*, **1995**, *51*, 2099. The hydroxy phosphonate is *O*-alkylated and then the protecting group is removed by treatment with either trifluoroacetic acid or tetrabutylammonium fluoride to generate the desired methoxy phosphonate analog.

Alternatively, the aldehyde is mixed with diethyl (2-aminoethyl)phosphonate and treated with a reducing agent such as sodium triacetoxyborohydride to generate the amino phosphonate analog.

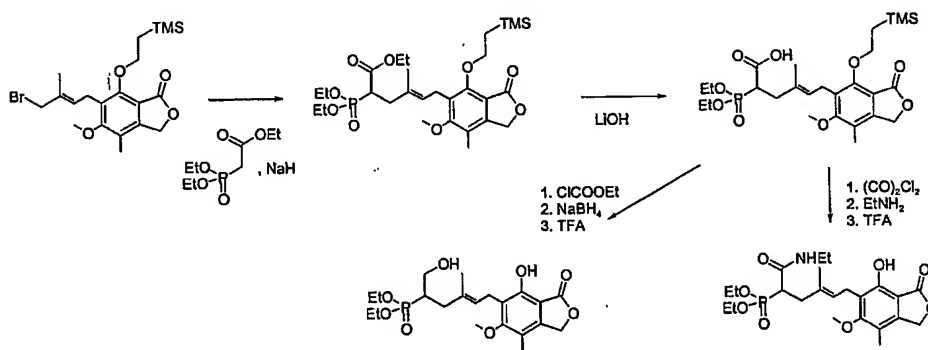


A solution of 4-[6-methoxy-7-methyl-3-oxo-4-(2-trimethylsilyl-ethoxy)-1,3-dihydro-isobenzofuran-5-yl]-2-methyl-but-2-enal (103 mg, 0.27 mmol) in methanol (5 mL) was cooled to 0°C. A solution of CeCl₃ (0.68 mL, MeOH: H₂O, 9:1) was added, followed by LiBH₄ (0.14 mL, 0.28 mmol of a 2M solution in THF). The ice bath was removed and the reaction mixture was allowed to warm to room temperature. The reaction mixture was stirred for an additional 40 minutes whereupon TLC indicated complete consumption of starting aldehyde. The reaction was worked up by addition of aqueous 1N HCl

(0.5 mL) and the product was extracted with CH_2Cl_2 . The organic layer was washed with saturated aqueous sodium bicarbonate solution and brine. The organic layer¹ was concentrated under reduced pressure and the residue was purified by silica gel chromatography to provide 100 mg (97%) of the product as a clear liquid. ^1H NMR (300 MHz, CDCl_3) δ 0.00 (s, 9H), 1.20 (dd, 2H, $J=7, 8$ Hz), 1.81 (s, 3H), 2.13 (s, 3H), 3.38-3.50 (m, 2H), 3.74 (s, 3H), 3.95 (s, 2H), 4.27 (dd, 2H, $J=7, 8$ Hz), 5.08 (s, 2H), 5.17-5.44 (m, 1H).

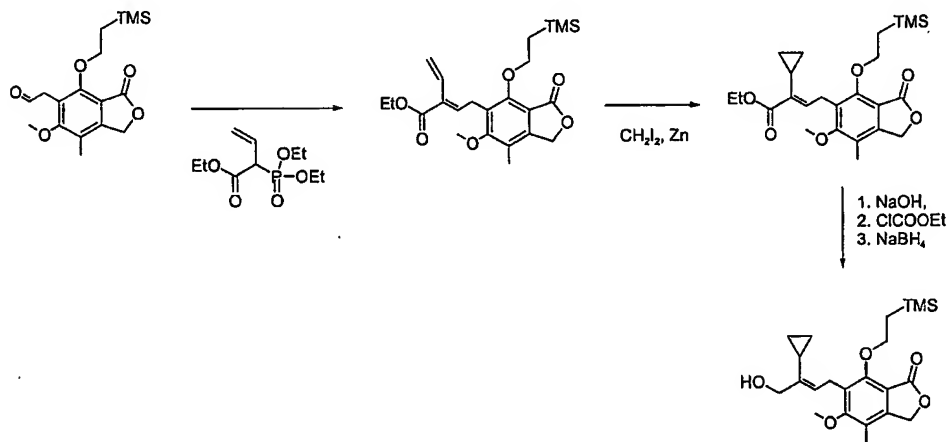


Polymer-supported triphenylphosphine is soaked in DCM for 1 hour. The allylic alcohol and carbon tetrabromide are sequentially added. When the reaction is complete, the mixture is filtered and the filtrate concentrated. The bromide is purified as necessary by chromatography.

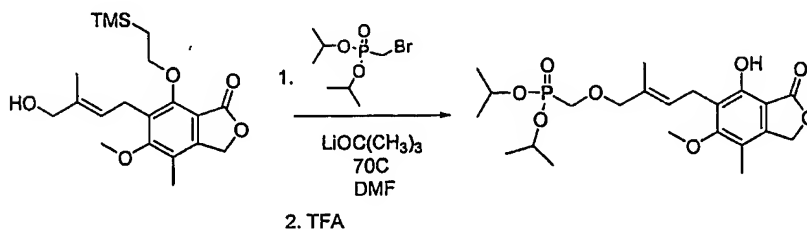


The allylic bromide is treated in an inert organic solvent such as dimethylformamide with an alkali metal salt of ethyl diethoxyphosphorylacetate (prepared by reacting ethyl diethoxyphosphorylacetate with sodium hexamethyldisilazide or sodium hydride) to afford the ethoxycarbonyl phosphonate, according to a procedure such as that described in WO 95/22538. The carboxylic ester group is converted to both the carboxylic amide and the hydroxymethyl groups according to the methods conventionally utilized for

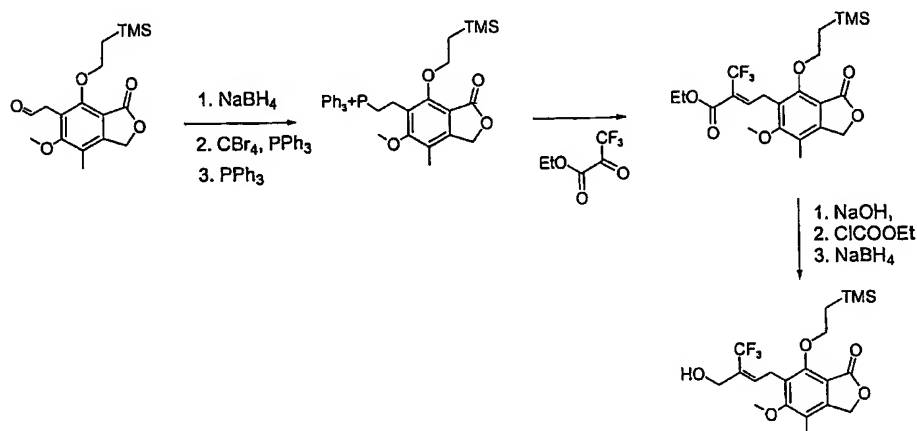
amide formations and ester reductions. For example, the carboxylic ester is saponified with aqueous lithium hydroxide. The acid is activated with ethyl chloroformate and reduced with sodium borohydride to generate, after removal of the protecting group, the hydroxymethyl phosphonate analog. The acid is also converted to its acyl chloride and then reacted with ethylamine to afford the amide analog.



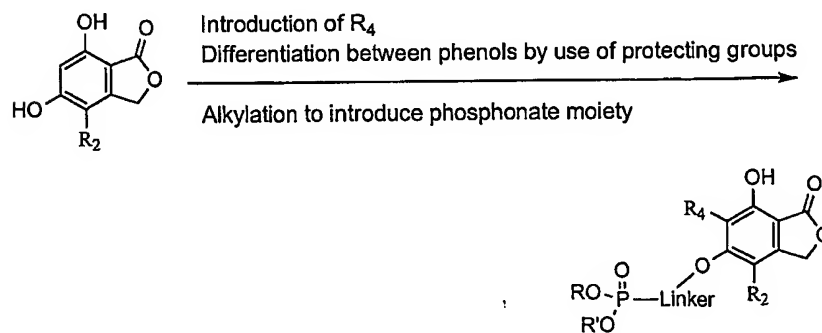
The aryl acetaldehyde is coupled with 2-(diethoxyphosphoryl)-but-3-enoic acid ethyl ester to generate the 2-vinyl substituted ester, according to a procedure such as that reported in *Synthesis*, 1999, 282. The 2-vinyl group is converted to the 2-cyclopropyl group under cyclopropanation conditions such as those described in *Tetrahedron Lett.* 1998, 39, 8621. The ester is converted to the alcohol, which, optionally, can be further subjected to reactions such as that described below to generate various phosphonate-containing mycophenolic acid analogues.



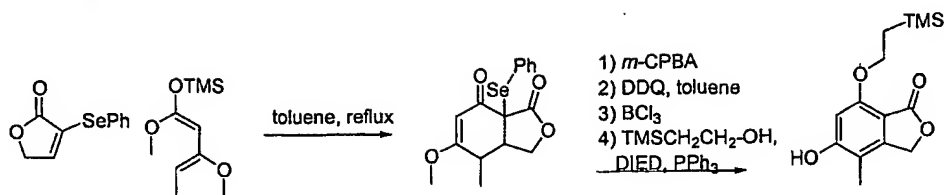
The allylic alcohol is treated with bromomethylphosphonic acid diisopropyl ester in the presence of a base such as lithium *t*-butoxide in a solvent such as dimethylformamide. The phenol protecting group is then removed by treatment with trifluoroacetic acid.



The phenacetaldehyde can alternatively be converted to the allyl phosphonium salt, according to a procedure such as that reported in *J. Org. Chem.* 1987, 52, 849. The phosphonium salt is then treated with the commercially available 3,3,3-trifluoro-2-oxo-propionic acid ethyl ester and a base such as sodium hydride to generate the 2-trifluoromethyl substituted ester. The ester is converted to the alcohol, which, optionally, can be further subjected to reactions described earlier to generate mycophenolic acid analogues with various side chains containing the phosphonate group.



Introduction of R₄ variants



The enone (synthesis reviewed in *Tetrahedron*, 1985, 41, 4881-4889) and the diene (*Chem. Pharm. Bull.*, 1989, 37, 2948-2951) are dissolved in an organic solvent such as toluene, stirred at room temperature for 24 hours and heated to reflux for additional 5 hours, according to a procedure from *J. Med. Chem.*, 1996, 39, 4181-4196. The reaction mixture is cooled to room temperature and the solvent removed *in vacuo*. The crude reaction product is further purified by chromatography.

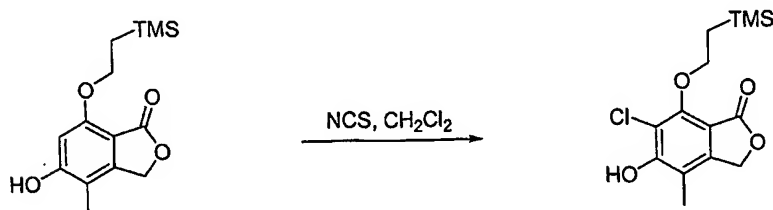
The product of step one is dissolved in an organic solvent such as DCM and *m*-chloroperbenzoic acid is added, according to a procedure from *J. Med. Chem.*, 1996, 39, 4181-4196. At the end of the reaction, the solution is poured into aqueous sodium hydrogen sulfite solution. The organic layer is washed with saturated aqueous sodium bicarbonate solution and is dried over sodium sulfate. Filtration and evaporation of solvents yields the crude product.

The crude product is dissolved in an organic solvent such as toluene and treated with dichlorodicyanoquinone (DDQ), according to a procedure from *J. Med. Chem.*, 1996, 39, 4181-4196. At the end of the reaction the solvent is removed *in vacuo* and the crude material is further purified by chromatography.

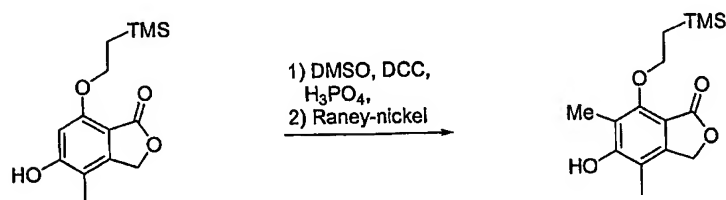
The product is dissolved in an organic solvent such as DCM and treated with boron trichloride at reflux temperature, according to a modified procedure from *J. Med. Chem.*, 1996, 39, 46-55. At the end of the reaction the solution is washed with aqueous HCl solution. The solution is dried over sodium sulfate. Removal of the solvent yields the crude reaction product. Further purification is achieved by chromatography.

The product of the previous step and triphenylphosphine are dissolved in an organic solvent such as tetrahydrofuran (THF). Diisopropylazodicarboxylate (DIAD) is added dropwise at 0°C. Stirring is continued. A solution of 2-trimethylsilyl ethanol in THF is added and stirring is continued. At the end of the reaction, the solvent is removed *in vacuo*. The crude reaction solid is

extracted with a mixture of organic solvents such as hexanes and diethylether. The washings are combined and the solvents removed *in vacuo*. The desired product is further purified and separated from the undesired regioisomer by chromatography.



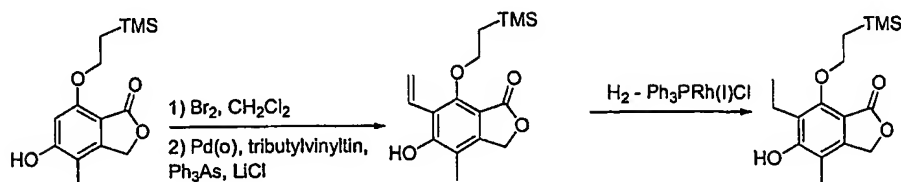
The starting material is dissolved in an organic solvent such as dimethylformamide (DMF) and reacted with *N*-chlorosuccinimide, according to a procedure from *J. Med. Chem.*, 1996, 39, 4181-4196. After the starting material is consumed the reaction mixture is poured into water and the product is extracted with diethyl ether. The combined organic layers are dried over sodium sulfate. Filtration and evaporation of the solvents yields the crude product. Further purification is achieved by chromatography.



The starting material is dissolved in an organic solvent such as benzene and reacted with dimethyl sulfoxide (DMSO), dicyclohexylcarbodiimide (DCC), and orthophosphoric acid according to a procedure from *J. Am. Chem. Soc.*, 1966, 88, 5855-5866. At the end of the reaction, the suspension is filtered and the organic layer washed with aqueous sodium bicarbonate solution and dried over sodium sulfate. Filtration and evaporation of solvents yields the crude material. Further purification is achieved by chromatography.

The product of step one is dissolved in an organic solvent such as DCM or THF and treated with Raney nickel, according to procedures reviewed in *Chem. Rev.*, 1962, 62, 347-404. When all starting material is consumed, the

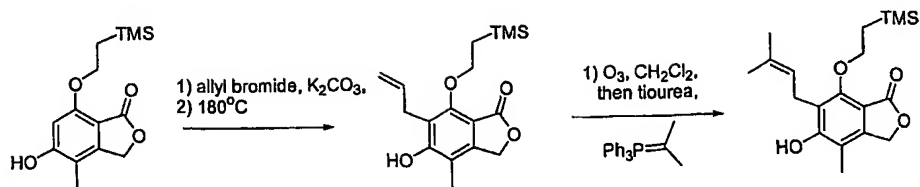
reaction is filtered and the solvent removed *in vacuo*. Further purification is achieved by chromatography.



The starting material is dissolved in an organic solvent such as DCM and bromine is added, according to a procedure from *J. Med. Chem.*, 1996, 39, 4181-4196. At the end of the reaction, additional DCM is added and the solution washed with aqueous sodium thiosulfate solution and brine. The organic layer is dried over sodium sulfate. Filtration and evaporation of solvents yields the crude material. Further purification is achieved by chromatography on silica gel.

The starting material, lithium chloride, triphenylarsine, tributylvinyltin, and tris(dibenzylideneacetone)dipalladium(0)-chloroform adduct are heated in an organic solvent such as *N*-methylpyrrolidinone at an elevated temperature of approximately 55°C, according to a procedure from *J. Med. Chem.*, 1996, 39, 4181-4196. At the end of the reaction, the mixture is cooled to room temperature and poured into a mixture of ice, potassium fluoride, water, and ethyl acetate. Stirring is continued for 1 hour. The suspension is filtered through Celite and extracted with ethyl acetate. The combined organic extracts are dried over sodium sulfate. The solvents are removed *in vacuo* and the crude material is further purified by chromatography.

The product of step two is dissolved in a mixture of organic solvents such as benzene and ethyl acetate. Tris(triphenylphosphine)rhodium(I) chloride is added and the reaction is placed under an atmosphere of hydrogen, according to a procedure from *J. Med. Chem.*, 1996, 39, 4181-4196. The solvents are removed *in vacuo* and the crude reaction is filtered through silica gel. Further purification is achieved by chromatography.

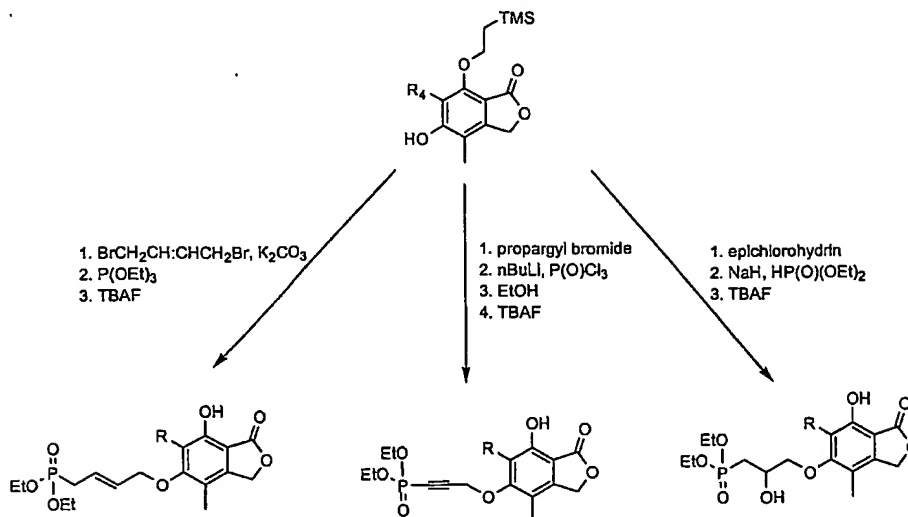


The starting material is dissolved in an organic solvent such as DMF. Potassium carbonate and allyl bromide are added and stirring at room temperature is continued, according to a procedure from *J. Med. Chem.*, **1996**, *39*, 4181-4196. After all the starting material is consumed, aqueous HCl solution and diethyl ether are added and the organic layer is collected and the solvent is removed *in vacuo*.

The crude material is dissolved in *N,N* diethylaniline and the reaction mixture is heated at an elevated temperature of *ca.* 180°C. At the end of the reaction, the mixture is cooled to room temperature and poured into a mixture of aqueous HCl (2N) and ethyl acetate. The organic layer is washed with aqueous HCl (2N) and dried over sodium sulfate. Filtration and removal of the solvents yields the crude product. Further purification is achieved by chromatography.

The product of step 2 is dissolved in a mixture of organic solvents such as methanol, DCM, and pyridine. The solution is cooled to -78°C and ozone is bubbled into the solution until a blue color persists. The excess ozone is removed with a nitrogen stream. The reaction mixture is warmed to room temperature and thiourea is added. Stirring at room temperature is continued. The reaction mixture is filtered and partitioned between DCM and water. The aqueous layer is extracted with DCM and the combined organic layers are washed with HCl (1 N), saturated aqueous sodium bicarbonate solution and brine. The solution is dried over sodium sulfate. Filtration and evaporation of the solvents yields the crude aldehyde. Further purification is achieved by chromatography.

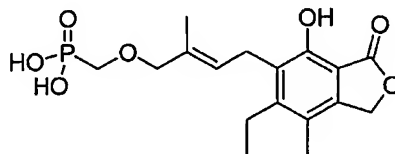
The aldehyde is dissolved in an organic solvent such as THF and is reacted with triphenylphosphonium *sec.*propyl bromide and potassium *tert.*butoxide, according to procedures reviewed in *Chem. Rev.*, **1989**, *89*, 863-927. At the end of the reaction, the solvent is removed *in vacuo* and the crude material purified by chromatography.

(A) Introduction of linkers to phosphonates

The phenols shown herein may optionally be alkylated with the reagent of choice. Optionally, the phosphonate moiety will be part of such a reagent. Alternatively, it will be introduced in a subsequent step by a variety of means, of which three are illustrated above. For example, an alkyl halide may be heated with triethylphosphite in a solvent such as toluene (or other Arbuzov reaction conditions: see Engel, R., "Synthesis of Carbon-phosphorus Bonds," CRC press, 1988). Alternatively, an epoxide may be reacted with the anion of a dialkyl phosphinate. In a further example, the phosphonate reagent may be the electrophile, *e.g.*, an acetylide anion may be condensed with phosphorus oxychloride and the intermediate dichlorophosphonate quenched with ethanol to generate the diethyl ester of the desired phosphonic acid.

Example 333

A specific compound of the invention can be prepared as follows.

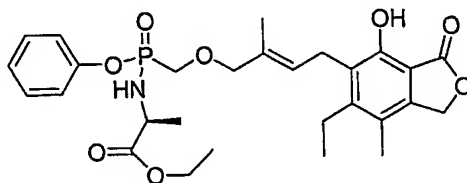


[4-(6-Ethyl-4-hydroxy-7-methyl-3-oxo-1,3-dihydro-isobenzofuran-5-yl)-2-methyl-but-2-enyloxymethyl]-phosphonic acid

This product was prepared using methods similar to those described herein, *e.g.*, in Examples 292 and 317. MS (negative mode): 369.3 [$M^+ - 1$].

Example 333A

A specific compound of the invention can be prepared as follows.

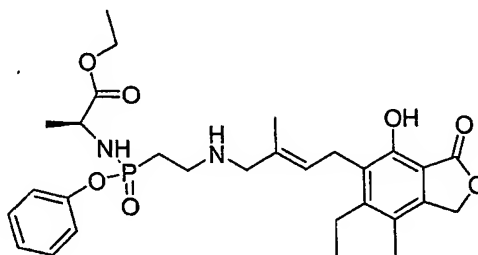


2-[[4-(6-Ethyl-4-hydroxy-7-methyl-3-oxo-1,3-dihydro-isobenzofuran-5-yl)-2-methyl-but-2-enyloxymethyl]-phenoxy-phosphinoylamino}-propionic acid ethyl ester

Using methods similar to those described herein, *e.g.*, in Example 302, the desired product was prepared, starting from Example 333. MS (positive mode): 546.3 [$M^+ + 1$] & 568.3 [$M^+ + Na$].

Example 334

A specific compound of the invention can be prepared as follows:

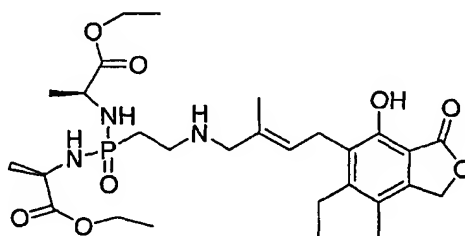


2-([2-[4-(6-Ethyl-4-hydroxy-7-methyl-3-oxo-1,3-dihydro-isobenzofuran-5-yl)-2-methyl-but-2-enylamino]-ethyl]-phenoxy-phosphinoylamino)-propionic acid ethyl ester

This product was prepared using methods analogous to those described herein, *e.g.*, in Examples 309 and 333, using 2-[(2-amino-ethyl)-phenoxy-phosphinoylamino]-propionic acid ethyl ester in the reductive amination step. MS (positive mode): 559.4 $[M^+ + 1]$ & 581.3 $[M^+ + Na]$.

Example 335

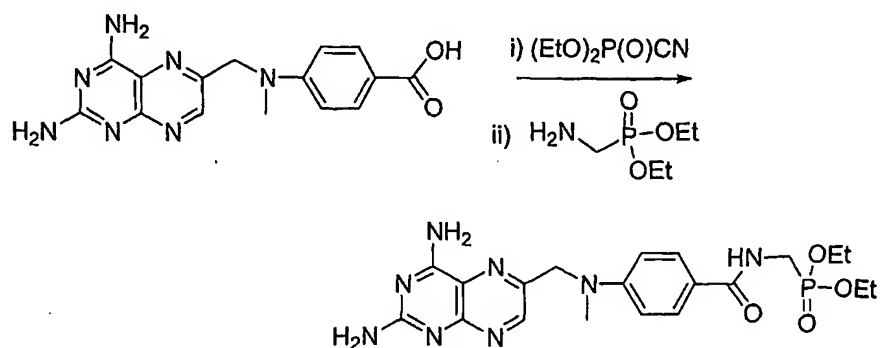
A specific compound of the invention can be prepared as follows:



2-((1-Ethoxycarbonyl-ethylamino)-{2-[4-(6-ethyl-4-hydroxy-7-methyl-3-oxo-1,3-dihydro-isobenzofuran-5-yl)-2-methyl-but-2-enylamino]-ethyl}-phosphinoylamino)-propionic acid ethyl ester

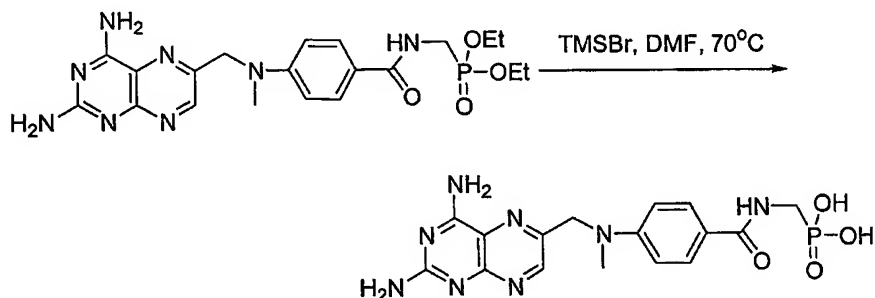
This product was prepared by methods analogous to those described herein, *e.g.*, in Example 334, using 2-[(2-aminoethyl)-(1-ethoxycarbonyl-ethylamino)-phosphinoylamino]-propionic acid ethyl ester in the reductive amination step. MS (positive mode): 582.4 $[M^+ + 1]$ & 604.3 $[M^+ + Na]$.

Example 336 (2-{4-[(2,4-Diamino-pteridin-6-ylmethyl)-methyl-amino]-benzoylamino}-methyl)-phosphonic acid diethyl ester



To a solution of 4-[(2,4-diamino-pteridin-6-ylmethyl)-methyl-amino]-benzoic acid hemihydrochloride dihydrate (67.0 mg, 177 μmol) in DMF (3.0 mL) was added diethyl cyanophosphonate (34.8 μL , 230 μmol) and diisopropylethylamine (Hunig's Base, DIEA, 30.4 μL , 177 μmol). The solution was stirred at ambient temperature for 4 hours when diethyl(aminomethyl)-phosphonate (45.4 mg, 177 μmol) was added. The solution was stirred for 4 additional hours, when complete consumption of the starting materials was observed. The reaction was worked up by removal of the solvent *in vacuo* and purifying the residue by silica gel chromatography using MeOH-CH₂Cl₂ (10-30%). The product collected from this chromatography step was sufficiently pure to be carried on to the next reaction. A small amount of the product (20 mg) was repurified by RP HPLC on C₁₈ column using H₂O/acetonitrile (2-95%) to provide 12.9 mg (76%) of the pure product. ¹H NMR (300 MHz, DMSO-*d*₆) δ 1.19 (t, 6H, *J* = 7.2 Hz), 3.21 (s, 3H), 3.70 (m, 2H), 4.00 (q, 4H, *J* = 7.2 Hz), 4.81 (s, 2H), 6.81 (d, 2H, *J* = 9 Hz), 7.71 (d, 2H, *J* = 9 Hz), 8.40 (br s, 1H), 8.61 (s, 1H). ³¹P (121.4 MHz, DMSO-*d*₆) δ 23.4. MS (*m/z*) 475.2 [M+H]⁺, 597.2 [M+Na]⁺.

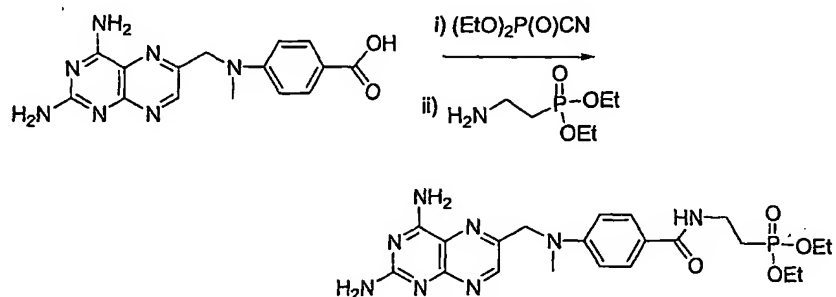
Example 337 (2-{4-[(2,4-Diamino-pteridin-6-ylmethyl)-methyl-amino]-benzoylamino}-methyl)-phosphonic acid



To a solution of crude (2-{4-[(2,4-Diamino-pteridin-6-ylmethyl)-methyl-amino]-benzoylamino}-ethyl)-phosphonic acid diethyl ester post silica column chromatography (60 mg, 126 μmol) in dry DMF (0.90 mL) was added trimethylsilyl bromide (bromotrimethylsilane, TMSBr, 130.6 μL , 1,010 μmol) at

ambient temperature. The solution was then heated at 70 °C for 4.0 hours, after which the reaction mixture was allowed to cool to room temperature. The solvent volume was reduced to ~ 700 μL *in vacuo* and diluted with H_2O (100 μL). This solution was purified by RP HPLC on C_{18} column using H_2O /acetonitrile (2-95%) to provide 26.8 mg (51%) of the desired compound as a yellow solid. ^1H NMR (300 MHz, $\text{DMSO}-d_6$) δ 3.18 (s, 3H), 3.50 (m, 2H), 4.77 (s, 2H), 6.79 (d, 2H, $J=9$ Hz), 7.79 (d, 2H, $J=9$ Hz), 8.07 (br s, 1H), 8.56 (s, 1H); MS (m/z) 419.2 $[\text{M}+\text{H}]^+$.

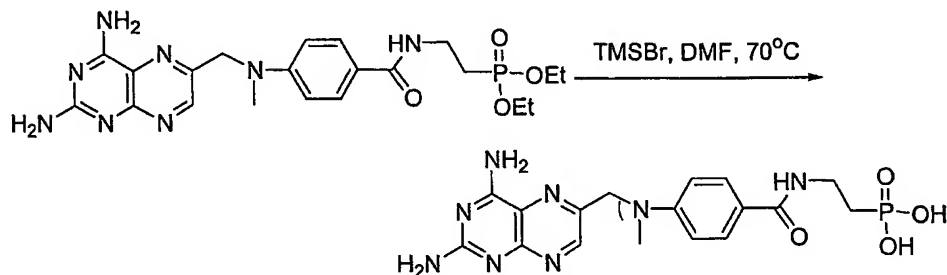
Example 338 (2-{4-[(2,4-Diamino-pteridin-6-ylmethyl)-methyl-amino]-benzoylamino}-ethyl)-phosphonic acid diethyl ester



To a solution of 4-[(2,4-diamino-pteridin-6-ylmethyl)-methyl-amino]-benzoic acid hemihydrochloride dihydrate (61.2 mg, 161 μmol) in DMF (2.8 mL) were added diethyl cyanophosphonate (31.8 μL , 210 μmol) and DIEA (27.8 μL , 161 μmol). The solution was stirred at ambient temperature for 4 hours, when diethyl(aminoethyl)phosphonate (43.8 mg, 161 μmol) was added. The solution was stirred for 3 additional hours, by which time complete consumption of the starting materials was observed. The reaction was worked up by removal of the solvent *in vacuo* and purifying the residue by silica gel chromatography using $\text{MeOH}-\text{CH}_2\text{Cl}_2$ (10-30%). The product collected from this chromatography step was sufficiently pure to be carried on to the next reaction. A small amount of the product (32 mg) was re-purified by RP HPLC on C_{18} column using H_2O /acetonitrile (2-95%) to provide 19 mg (70%) of the pure product. ^1H NMR (300 MHz, $\text{DMSO}-d_6$) δ 1.21 (t, 6H, $J=7$ Hz), 1.95- 2.05 (m,

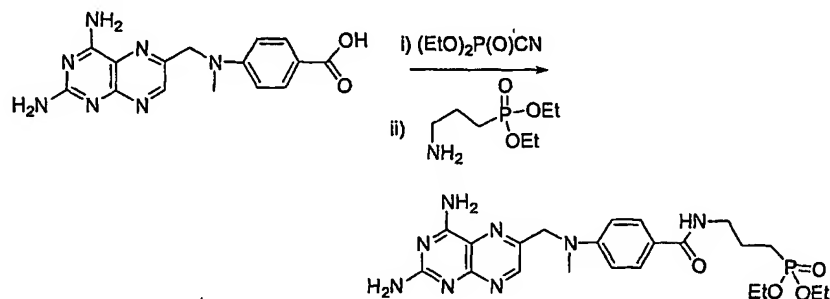
2H), 3.20 (s, 3H), 3.13- 3.22 (m, 2H), 3.98 (appt septet, 4H, $J=7$ Hz), 4.79 (s, 2H), 6.80 (d, 2H, $J=9$ Hz), 7.65 (d, 2H, $J=9$ Hz), 8.20 (br s, 1H), 8.60 (s, 1H). ^{31}P (121.4 MHz, $\text{DMSO}-d_6$) δ 28.9. MS (m/z) 489.2 $[\text{M}+\text{H}]^+$, 511.2 $[\text{M}+\text{Na}]^+$.

Example 339 (2-{4-[(2,4-Diamino-pteridin-6-ylmethyl)-methyl-amino]-benzoylamino}-ethyl)-phosphonic acid



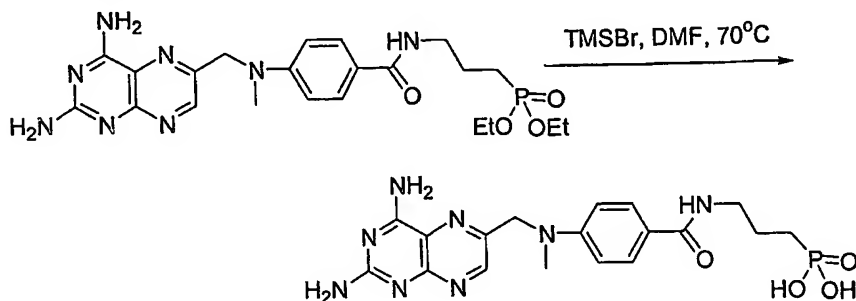
To a solution of crude (2-{4-[(2,4-diamino-pteridin-6-ylmethyl)-methyl-amino]-benzoylamino}-ethyl)-phosphonic acid diethyl ester post silica column chromatography (61 mg, 125 μmol) in dry DMF (1.00 mL) was added TMSBr (129.0 μL , 999.2 μmol) at ambient temperature. The solution was then heated at 70 °C for 5.5 hours, when LCMS analysis demonstrated the reaction to be 90% complete. The reaction mixture was allowed to cool to room temperature and stirred for an additional 12 hours. The reaction was worked up by removal of the solvent *in vacuo* and dissolving the residue in DMF / H_2O (800 μL , 1:1) and 1N aqueous NaOH (15 μL). The product was purified by RP HPLC on C_{18} column using H_2O /acetonitrile (2-95%) to provide 29 mg (53%) of the desired compound as a yellow solid. ^1H NMR (300 MHz, $\text{DMSO}-d_6$) δ 1.67- 1.85 (m, 2H), 3.19 (s, 3H), 3.25- 3.40 (m, 2H), 4.76 (s, 2H), 6.71 (br s, 2H), 5.80 (d, 2H, $J=9$ Hz), 7.64 (d, 2H, $J=9$ Hz), 7.73 (br s, 2H), 8.15 (br s, 1H), 8.56 (s, 1H). ^{31}P (121.4 MHz, $\text{DMSO}-d_6$) δ 23.0. MS (m/z) 431.3 $[\text{M}-\text{H}]^-$.

Example 340 (2-{4-[(2,4-Diamino-pteridin-6-ylmethyl)-methyl-amino]-benzoylamino}-propyl)-phosphonic acid diethyl ester



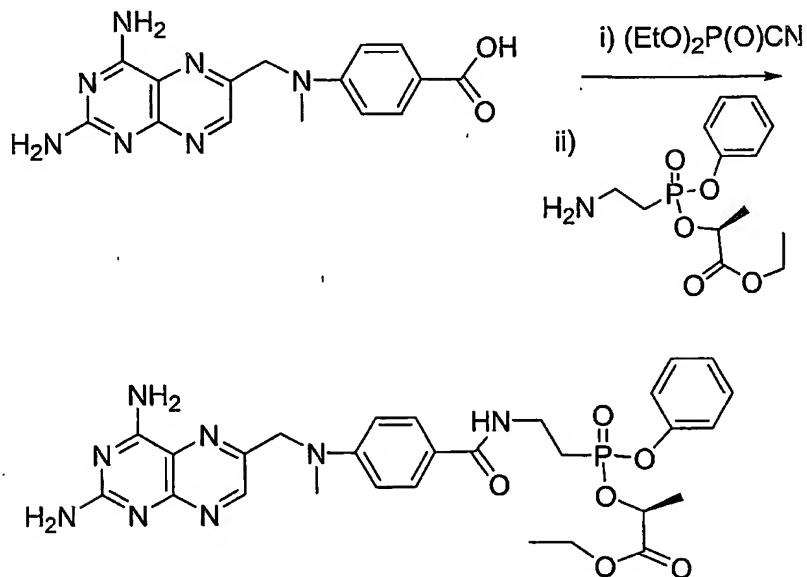
To a solution of 4-[(2,4-diamino-pteridin-6-ylmethyl)-methyl-amino]-benzoic acid hemihydrochloride dihydrate (61.2 mg, 161 μmol) in DMF (2.8 mL) were added diethyl cyanophosphonate (31.8 μL , 210 μmol) and DIEA (27.8 μL , 161 μmol). The solution was stirred at ambient temperature for 3 hours, when diethyl(aminopropyl)phosphonate (34.9 mg, 122.6 μmol) was added. The solution was stirred for 2 additional hours, whereupon complete consumption of the starting materials was observed. The reaction was worked up by removal of the solvent *in vacuo* and purifying the residue by silica gel chromatography using $\text{MeOH}-\text{CH}_2\text{Cl}_2$ (10-30%). The product (65.5 mg) collected from this chromatography step was sufficiently pure to be carried on to the next reaction. A small amount (32.8 mg) was re-purified by RP HPLC on C_{18} column using H_2O /acetonitrile (2-95%) to provide 23.2 mg (75%) of the pure product. ^1H NMR (300 MHz, $\text{DMSO}-d_6$) δ 1.20 (t, 6H, $J=7.2$ Hz), 1.64-1.75 (m, 4H), 3.22 (s, 3H), 3.41 (m, 2H), 3.98 (appt septet, 4H, $J=7.2$ Hz), 4.85 (s, 2H), 6.79 (d, 2H, $J=9$ Hz), 7.68 (d, 2H, $J=9$ Hz), 8.17 (br s, 1H), 8.70 (s, 1H); ^{31}P (121.4 MHz, $\text{DMSO}-d_6$) δ 31.9; MS (m/z) 503.2 $[\text{M}+\text{H}]^+$.

Example 341 (2-{4-[(2,4-Diamino-pteridin-6-ylmethyl)-methyl-amino]-benzoylamino}-propyl)-phosphonic acid



To a solution of crude (2-{4-[(2,4-diamino-pteridin-6-ylmethyl)-methyl-amino]-benzoylamino}-propyl)-phosphonic acid diethyl ester post silica column chromatography (32.2 mg, 66.2 μmol) in dry DMF (0.50 mL) was added TMSBr (68.0 μL , 529.6 μmol) at ambient temperature. The solution was then heated at 70 °C for 1.0 hour, when LCMS analysis demonstrated the reaction to be complete. The reaction mixture was allowed to cool to room temperature, and water (60 μL) and methanol (60 μL) were added. The crude reaction mixture was purified by RP HPLC on C₁₈ column using H₂O/acetonitrile (2-95%) to provide 11.2 mg (38%) of the desired compound as a yellow solid. ¹H NMR (300 MHz, DMSO-*d*₆) δ 1.50 (m, 2H), 1.61 (m, 2H), 3.22 (s, 3H), 3.25- 3.40 (m, 2H), 4.84 (s, 2H), 6.80 (d, 2H, *J*= 9 Hz), 7.69 (d, 2H, *J*= 9 Hz), 8.20 (br s, 1H), 8.69 (s, 1H). ³¹P (121.4 MHz, DMSO-*d*₆) δ 26.3. MS (*m/z*) 447.3 [M-H]⁻.

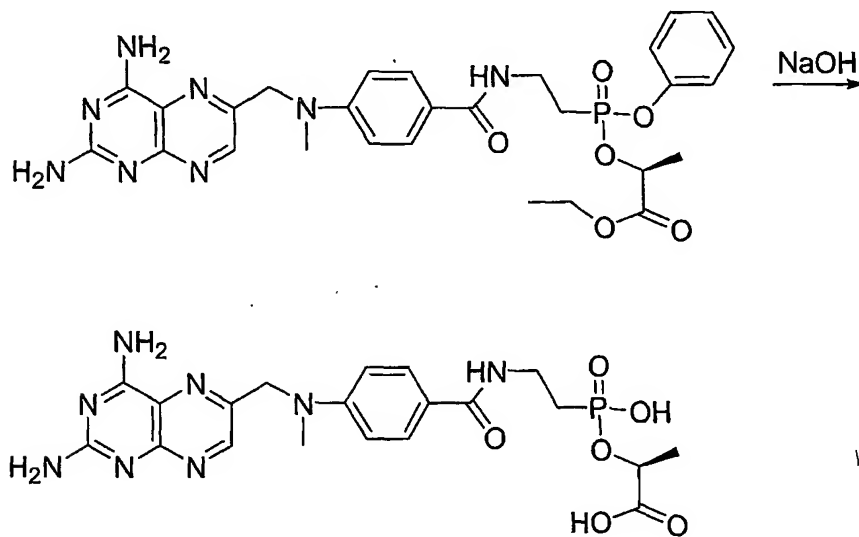
Example 342 2-[(2-{4-[(2,4-diaminopteridin-6-ylmethyl)methyl-amino]benzoylamino}-ethyl)phenoxyphosphinoyloxy]propionic acid ethyl ester [diastereomeric mixture at phosphorus]



To a solution of 4-[(2,4-diamino-pteridin-6-ylmethyl)-methyl-amino]-benzoic acid hemihydrochloride dihydrate (60.0 mg, 158.3 μmol) in DMF (2.5 mL) were added diethyl cyanophosphonate (31.2 μL , 205.7 μmol) and DIEA (81.8 μL , 474.9 μmol). The solution was stirred at ambient temperature for 3.5 hours, when a solution of (S)-2-[(2-aminoethyl)phenoxyphosphinoyloxy]-propionic acid ethyl ester mono acetic acid salt (57.1 mg, 158.3 μmol ; mixture of diastereomers at phosphorus) in DMF (200 μL) was added. The solution was stirred for 1.5 additional hours, whereupon complete consumption of the starting materials was observed. The solvent was removed *in vacuo* and the crude material was purified by silica gel chromatography using MeOH-CH₂Cl₂ (10-30%). A small amount of the product (24.8 mg) was repurified by RP HPLC on C₁₈ column using H₂O/acetonitrile (2-95%) to provide 15.8 mg (65%) of the pure product. ¹H NMR (300 MHz, DMSO-*d*₆) δ 1.17 – 1.27 (m, 3H), 1.32 (d, 2H, J = 7.5 Hz), 1.42 (d, 1H, J = 7.5 Hz), 2.27 (m, 2H), 3.19 (s, 3H), 3.53 (m, 2H), 4.08 – 4.14 (m, 2H), 4.77 (s, 2H), 4.98 (m, 1H), 6.72 (br s, 1H), 6.81 (d,

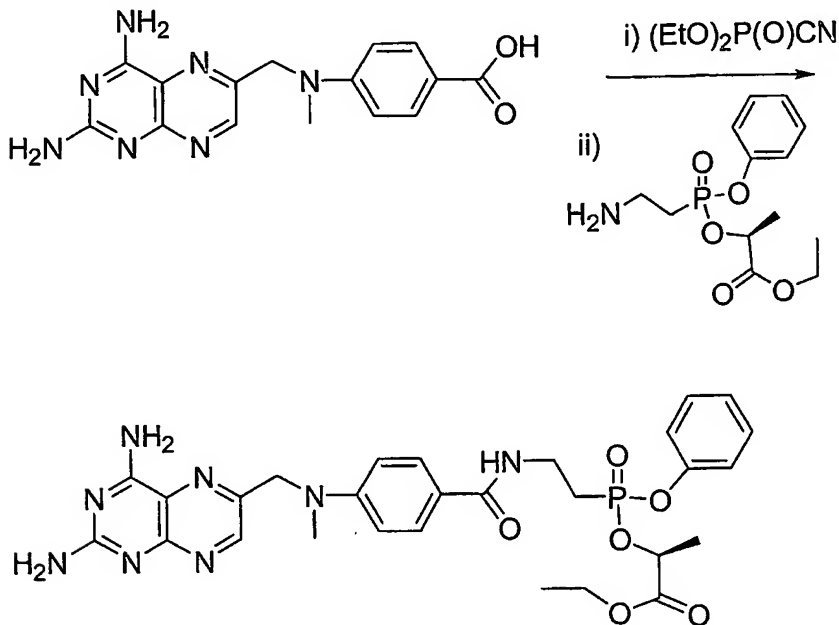
2H, $J = 9$ Hz), 7.21 (m, 3H), 7.36 (m, 2H), 7.66 (d, 2H, $J = 9$ Hz), 8.26 (br s, 1H), 8.56 (s, 1H); ^{31}P (121.4 MHz, DMSO- d_6) δ 26.6, 27.4. MS (m/z) 609.2 $[\text{M}+\text{H}]^+$.

Example 343 2-[(2-{4-[(2,4-diaminopteridin-6-ylmethyl)methylamino]benzoylamino}-ethyl)phenoxyphosphinoyloxy]-propionic acid [diastereomeric mixture at phosphorus]



To a solution of 2-[(2-{4-[(2,4-diaminopteridin-6-ylmethyl)methylamino]benzoylamino}ethyl)phenoxy-phosphinoyloxy]propionic acid ethyl ester (mixture of diastereomers at phosphorus; 40.0 mg, 65.7 μmol) in DMF (0.4 mL), acetonitrile (0.2 mL) and water (0.2 mL) was added aqueous sodium hydroxide (1 N, 131.4 μL). The solution was stirred at ambient temperature for 4 hours. The solvents were removed *in vacuo* and the crude product was purified by RP HPLC on C₁₈ column using H₂O/acetonitrile (2-95%) to provide 23.7 mg (71.3%) of the pure product. ^1H NMR (300 MHz, DMSO- d_6) δ 1.30 (d, 2H, $J = 6.9$ Hz), 1.79 (m, 2H), 3.21 (s, 3H), 3.37 (m, 2H), 4.61 (m, 1H), 4.81 (s, 2H), 6.79 (d, 2H, $J = 8.7$ Hz), 7.64 (d, 2H, $J = 9.7$ Hz), 8.25 (br s, 1H), 8.63 (s, 1H); ^{31}P (121.4 MHz, DMSO- d_6) δ 25.1. MS (m/z) 505.2 $[\text{M}+\text{H}]^+$.

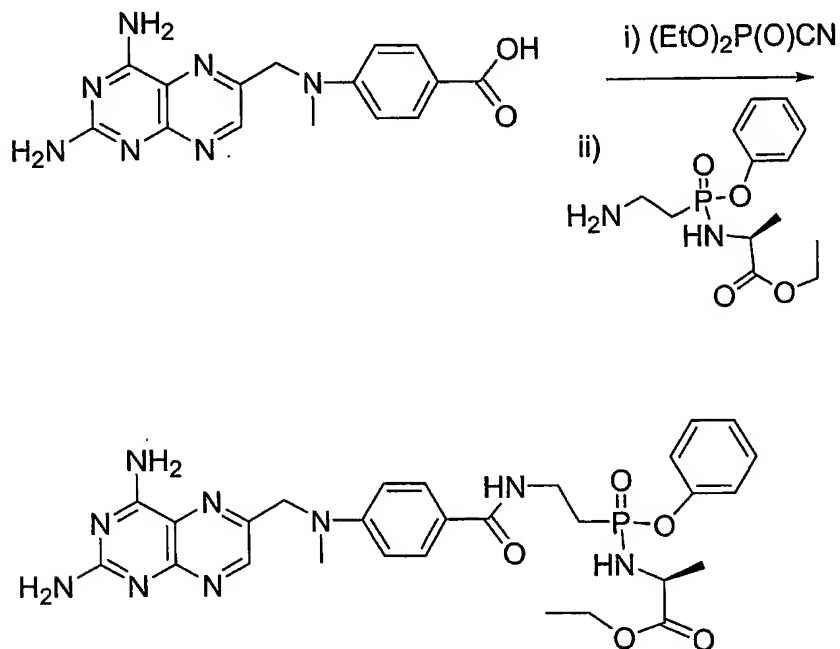
Example 343A 2-[(2-{4-[(2,4-diaminopteridin-6-ylmethyl)methylamino]-benzoylamino}ethyl)phenoxyphosphinoyloxy]propionic acid ethyl ester [diastereomerically pure at phosphorus]



To a solution of 4-[(2,4-diaminopteridin-6-ylmethyl)-methyl-amino]benzoic acid hemihydrochloride dihydrate (101.9 mg, 268.9 μmol) in DMF (3.3 mL) were added diethyl cyanophosphonate (53.0 μL , 349.5 μmol) and DIEA (138.0 μL , 806.7 μmol). The solution was stirred at ambient temperature for 2.5 hours, whereupon (S)-2-[(2-aminoethyl)phenoxyphosphinoyloxy]-propionic acid ethyl ester mono acetic acid salt (diastereomerically pure at phosphorus; 268.9 μmol) in DMF (500 μL) was added. The solution was stirred for 30 additional minutes, whereupon complete consumption of the starting materials was observed. The solvent was removed *in vacuo* and the crude material was purified by silica gel chromatography using MeOH-CH₂Cl₂ (10-30%). A small amount of the product (40.0 mg) was repurified by RP HPLC on C₁₈ column using H₂O/acetonitrile (2-95%) to provide 28.7 mg (75.1%) of the pure product. ¹H NMR (300 MHz, DMSO-*d*₆) δ 1.15 (t, 3H, J = 7.2 Hz), 1.44 (d, 3H, J = 6.9 Hz), 2.26 (m, 2H), 3.23 (s, 3H), 3.51 (m, 2H), 4.09 (q, 2H, J = 7.2 Hz), 4.86 (s, 2H), 5.01 (m, 1H), 6.81 (d, 2H, J = 9.3 Hz), 7.21 (m, 3H), 7.35 (m,

2H), 7.68 (d, 2H, $J = 9.3$ Hz), 8.29 (br s, 1H), 8.71 (s, 1H); ^{31}P (121.4 MHz, DMSO- d_6) δ 26.6. MS (m/z) 609.2 $[\text{M}+\text{H}]^+$.

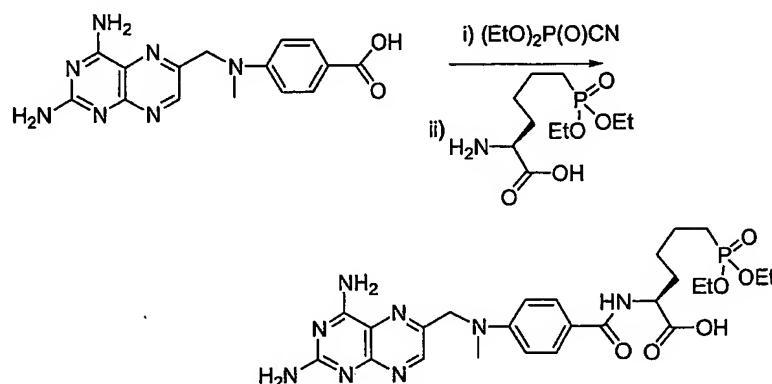
Example 344 2-[(2-{4-[(2,4-diaminopteridin-6-ylmethyl)methylamino]-benzoylamino}-ethyl)-phenoxyphosphinoylamino]propionic acid ethyl ester (mixture of diastereomers at phosphorus)



To a solution of 4-[(2,4-diamino-pteridin-6-ylmethyl)-methyl-amino]-benzoic acid hemihydrochloride dihydrate (39.6 mg, 104.0 μmol) in DMF (1.2 mL) were added diethyl cyanophosphonate (20.6 μL , 136.1 μmol) and DIEA (36.0 μL , 209.4 μmol). The solution was stirred at ambient temperature for 3 hours, when (S)-2-[(2-aminoethyl)phenoxyphosphinoylamino]propionic acid ethyl ester mono acetic acid salt (mixture of diastereomers at phosphorus; 104.0 μmol) in DMF (200 μL) was added. The solution was stirred for 30 minutes when complete consumption of the starting materials was observed. An aliquot (66%) of the reaction was purified by silica gel chromatography using MeOH- CH_2Cl_2 (10-30%), yielding 27.2 mg of crude product. A small amount of the product (10 mg) was repurified by RP HPLC on C_{18} column using

H₂O/acetonitrile (2-95%) to provide 4.2 mg (26%) of the pure product. ¹H NMR (300 MHz, DMSO-*d*₆) δ 1.11 (t, 3H, *J* = 6.9 Hz), 1.18 (d, 3H, *J* = 7.2 Hz), 2.06-2.17 (m, 2H), 3.20 (s, 3H), 3.51 (m, 2H), 3.88 (m, 1H), 4.02 (m, 2H), 4.79 (s, 2H), 5.61 (m, 1H), 6.80 (d, 2H, *J* = 9 Hz), 6.98 (br s, 1H); 7.18 (m, 3H), 7.32 (m, 2H), 7.67 (d, 2H, *J* = 9 Hz), 8.20 (br s, 1H), 8.59 (s, 1H). ³¹P (121.4 MHz, DMSO-*d*₆) δ 29.5, 30.1. MS (*m/z*) 608.2 [M+H]⁺.

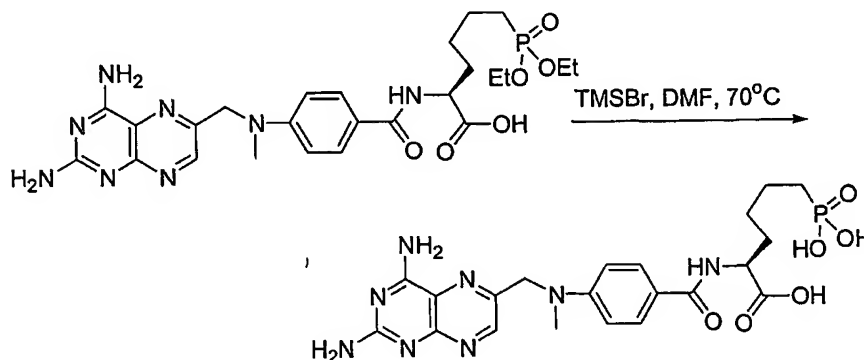
Example 345 2-{4-[(2,4-Diamino-pteridin-6-ylmethyl)-methyl-amino]-benzoylamino}-6-(diethoxy-phosphoryl)-hexanoic acid



To a solution of 4-[(2,4-diamino-pteridin-6-ylmethyl)-methyl-amino]-benzoic acid hemihydrochloride dihydrate (63.0 mg, 166.2 μmol) in DMF (2.8 mL) were added diethyl cyano phosphonate (30.8 μL, 199.4 μmol) and DIEA (85.8 μL, 498.6 μmol). The solution was stirred at ambient temperature for 3.5 hours when (L)-2-amino-6-diethylphosphonohexanoic acid (44.3 mg, 166.2 μmol) was added. The solution was stirred for 48 additional hours. The reaction was worked up by removal of the solvent *in vacuo* and purifying the residue by silica gel chromatography using MeOH-CH₂Cl₂ (10-30%). The product (87 mg) collected from this chromatography step was sufficiently pure to be carried on to the next reaction. An aliquot of the product (51.0 mg) was repurified by RP HPLC on C₁₈ column using H₂O/acetonitrile (2-95%) to provide 24.7 mg (44%) of the pure product. ¹H NMR (300 MHz, DMSO-*d*₆) δ 1.18 (t, 6H, *J* = 6.9 Hz), 1.42 (m, 4H), 1.65 (m, 4H), 3.20 (s, 3H), 3.92 (m, 4H), 4.29 (m, 1H), 4.78 (s, 2H), 6.72 (br s, 1H), 6.81 (d, 2H, *J* = 9 Hz), 7.73 (d, 2H, *J* = 9 Hz), 8.14 (d, 1H, *J*

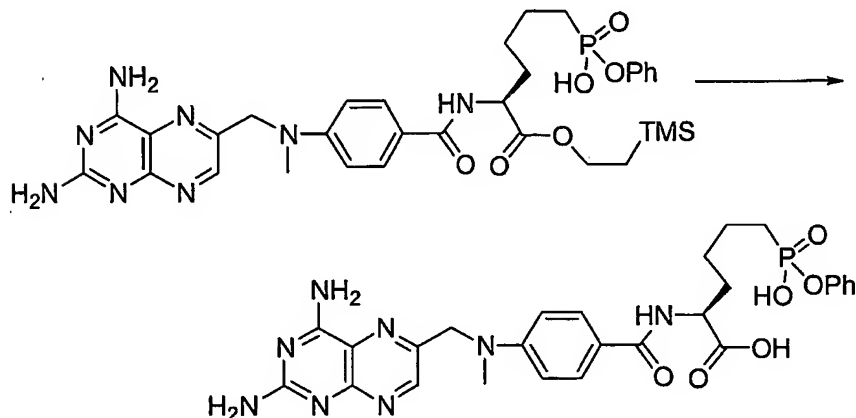
= 7.8 Hz), 8.56 (s, 1H); ^{31}P (121.4 MHz, $\text{DMSO-}d_6$) δ 31.8; MS (m/z) 574.3 $[\text{M}]^+$.

Example 346 2-{4-[(2,4-Diaminopteridin-6-ylmethyl)methylamino]-benzoylamino}-6-(phosphoryl)hexanoic acid



To a solution of crude (2-{4-[(2,4-diamino-pteridin-6-ylmethyl)-methylamino]-benzoylamino})-(L)-(6'-(phosphonic acid diethyl ester) hexanoic acid) post silica column chromatography (20 mg, 34.6 μmol) in dry DMF (0.60 mL) was added TMSBr (18.0 μL , 139.2 μmol) at ambient temperature. The solution was then heated at 70 °C for 18 hours, after which the reaction mixture was allowed to cool to room temperature. The solvent was removed *in vacuo* and dissolved in DMF (400 μL) and water (60 μL). This solution was purified by RP HPLC on C_{18} column using H_2O /acetonitrile (2-95%) to provide 8.9 mg (49%) of the product as a yellow solid. ^1H NMR (300 MHz, $\text{DMSO-}d_6$) δ 1.45 (m, 6H), 1.75 (m, 2H), 3.20 (s, 3H), 4.25 (m, 1H), 4.77 (s, 2H), 6.62 (br s, 1H), 6.80 (d, 2H, $J=8.7$ Hz), 7.73 (d, 2H, $J=8.7$ Hz), 8.14 (br s, 1H), 8.55 (s, 1H); MS (m/z) 519.2 $[\text{M}+\text{H}]^+$.

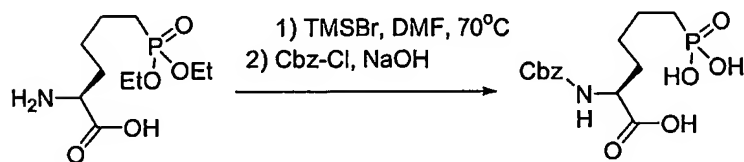
Example 347 2-{4-[(2,4-Diamino-pteridin-6-ylmethyl)-methyl-amino]-benzoylamino}-6'-(mono phenyl-phosphonate)hexanoic acid



The ethyl-TMS ester is hydrolyzed under suitable conditions to provide the corresponding acid of the invention.

The intermediate 2-{4-[(2,4-Diamino-pteridin-6-ylmethyl)-methyl-amino]-benzoylamino}-6'-(mono phenyl-phosphonate)-hexanoic acid TMS ethanol ester can be prepared as follows.

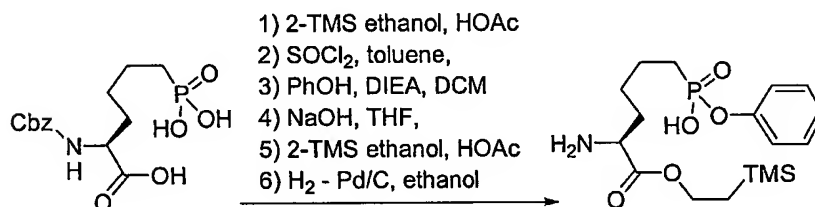
a. (L)-2- Cbz-Amino-hexanoic acid –6-phosphonic acid.



To a suspension of (L)-2-amino-6-(diethoxyphosphonyl)hexanoic acid (106 mg, 396.8 μ mol) in dry DMF (2.00 mL) was added TMSBr (307.0 μ L, 2,381.0 μ mol) at ambient temperature. The solution was then heated at 70 °C for 2 hours, after which the reaction mixture was allowed to cool to room temperature. The solvent was removed *in vacuo*. The crude material was

dissolved in water (0.25 mL) and NaOH (1-N, 2.50 mL). Benzyl chloroformate (79.3 μ L, 555.5 μ mol) was added and stirring at room temperature was continued. After 2 hours, the solution was washed with ether (2 mL) and the aqueous layer was acidified with aqueous HCl to pH 1. The aqueous layer was extracted with EtOAc (3x 5 mL). The combined organic extracts were dried over sodium sulfate. Filtration and evaporation of solvents yielded a crude product, which was sufficiently pure for further transformations. ^1H NMR (300 MHz, DMSO- d_6) δ 1.42 – 1.65 (m, 8H), 3.90 (m, 1H), 5.02 (s, 2H), 7.32 (s, 5H), 7.55 (m, 1H), 7.94 (s, 1H); ^{31}P (121.4 MHz, DMSO- d_6) δ 26.5; MS (m/z) 345.6 $[\text{M}+\text{H}]^+$.

b. (L)-2-Amino-hexanoic acid 2' TMS ethyl ester-6-phosphonic acid mono phenyl ester



To a solution of (L)-2-Cbz-amino-hexanoic acid-6-phosphonic acid (137.3 mg, 397.9 μ mol) in 2-TMS ethanol (2.5 mL) was added acetyl chloride (50 μ L). Stirring at room temperature was continued. After 22 hours complete conversion was observed. The solvents were removed *in vacuo*. The crude material was sufficiently pure for the next step.

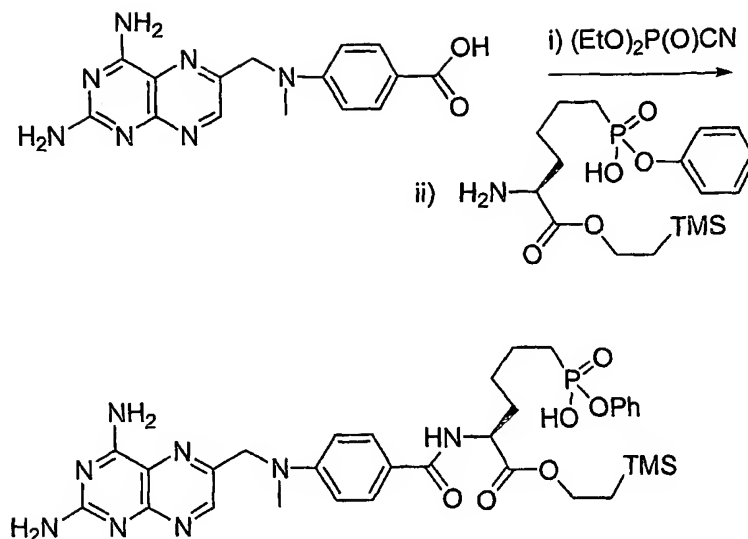
One half of the crude material (198.9 μ mol) was dissolved in toluene (3.0 mL) at room temperature. Thionyl chloride (167.2 mg, 1,416.0 μ mol) was added and the reaction mixture was heated at 70 $^\circ\text{C}$ (oil bath). After 4 hours, the reaction was cooled to room temperature and the solvent was removed *in vacuo*. The crude material was re-dissolved in methylene chloride (2.0 mL) and a solution of phenol (36.6 mg, 389.0 μ mol) and DIEA (67.0 μ L, 389.0 μ mol) in

methylene chloride (1.0 mL) was added. Stirring at room temperature was continued. After 4 hrs the solvents were removed *in vacuo*.

The crude material was dissolved in tetrahydrofuran (THF) (3.0 mL) and aqueous sodium hydroxide solution (1N, 0.885 mL) was added. Stirring at room temperature was continued. After 14 hours the solvent was removed *in vacuo* to provide the crude phosphonate mono phenyl ester (63.8 mg). This material was dissolved in 2-TMS ethanol (1.0 mL) and acetyl chloride (20 μ L) was added. Stirring at room temperature was continued. After 22 hours complete conversion to the carboxylate ester was observed. The solvents were removed *in vacuo*. The material was sufficiently pure for the next step.

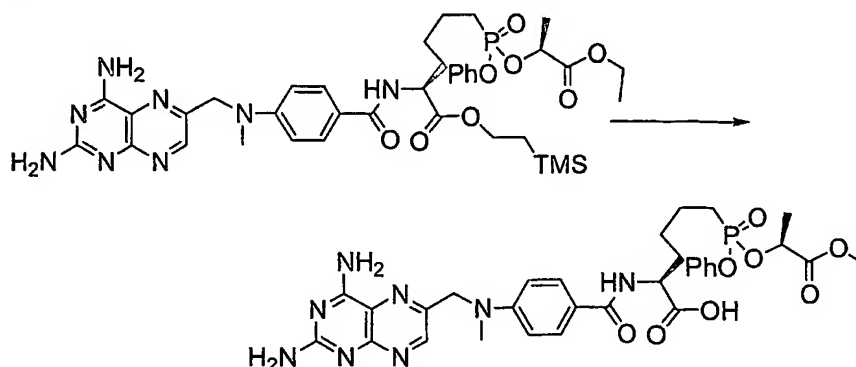
One half of the crude material (75 μ mol) was dissolved in ethanol (1.5 mL). Pd/C (5%, 20 mg) was added and the reaction was placed under an atmosphere of hydrogen gas. After 1.5 hours Celite was added and the crude reaction mixture was filtered through Celite. The solvents were removed *in vacuo* and the crude material was used in the next step without further purification.

c. 2-{4-[(2,4-diamino-pteridin-6-ylmethyl)-methyl-amino]-benzoylamino}-6'-(mono phenyl-phosphonate)-hexanoic acid TMS ethanol ester



To a solution of 4-[(2,4-diamino-pteridin-6-ylmethyl)-methyl-amino]-benzoic acid hemihydrochloride dihydrate (22.7 mg, 60.0 μmol) in DMF (0.80 mL) were added diethyl cyano phosphonate (12.4 μL , 78.0 μmol) and DIEA (31.0 μL , 180.0 μmol). The solution was stirred at ambient temperature for one hour when (L)-2-amino-6-monophenoxyposphonatohexanoic acid 2' TMS ethyl ester (70.5 μmol), suspended in DMF (0.2 mL), was added. The solution was stirred for 3.5 additional hours. The crude reaction mixture was purified by RP HPLC on C_{18} column using H_2O /acetonitrile (5-95%) to provide 19.4 mg (46%) of 2-{4-[(2,4-diamino-pteridin-6-ylmethyl)-methyl-amino]-benzoylamino}-6'-(mono phenyl-phosphonate)-hexanoic acid TMS ethanol ester. ^1H NMR (300 MHz, $\text{DMSO}-d_6$) δ 0.0 (s, 9H), 0.91 (t, 2H, $J=8.1$ Hz), 1.42 – 1.53 (m, 4H), 1.67 – 1.76 (m, 4H), 3.24 (s, 3H), 4.10 (t, 2H, $J=8.1$ Hz), 4.29 (m, 1H), 4.86 (s, 2H), 6.81 (d, 2H, $J=9$ Hz), 7.12 (m, 3H), 7.31 (m, 2H), 7.74 (d, 2H, $J=9$ Hz), 8.14 (d, 1H, $J=7.8$ Hz), 8.71 (s, 1H); ^{31}P (121.4 MHz, $\text{DMSO}-d_6$) δ 26.2; MS (m/z) 695.2 $[\text{M}]^+$.

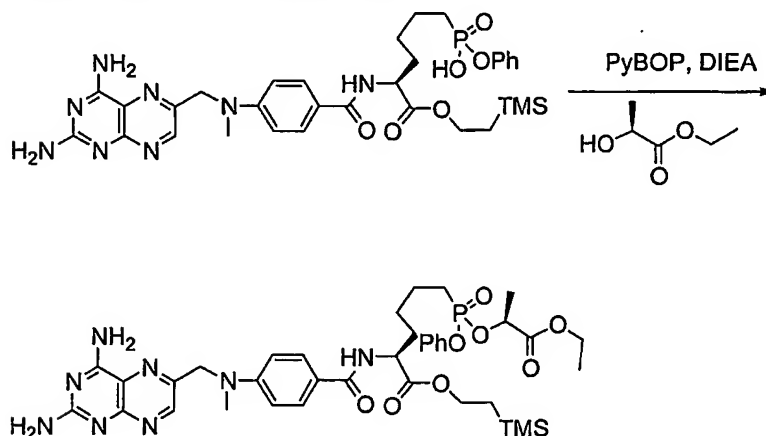
Example 347A 4-[(2,4-Diamino-pteridin-6-ylmethyl)methylamino]-benzoylamino}-6'--(mono phenyl mono (S) ethyl lactate-phosphonate)hexanoic acid



The ethyl-TMS ester is hydrolyzed under suitable conditions to provide the corresponding acid of the invention.

The intermediate 2-{4-[(2,4-diamino-pteridin-6-ylmethyl)-methyl-amino]-benzoylamino}-6'-1-(mono phenyl mono (S) ethyl lactate-phosphonate)-hexanoic acid TMS ethanol ester can be prepared as follows.

a. 2-{4-[(2,4-Diamino-pteridin-6-ylmethyl)-methyl-amino]-benzoylamino}-6'-1-(mono phenyl mono (S) ethyl lactate-phosphonate)-hexanoic acid TMS ethanol ester

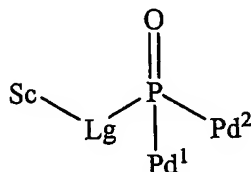


To a solution of 2-{4-[(2,4-diamino-pteridin-6-ylmethyl)-methyl-amino]-benzoylamino}-6'-1-(mono phenyl-phosphonate)-hexanoic acid TMS ethanol ester

(14.5 mg, 20.8 μ mol, Example 225) in DMF (0.70 mL) was added PyBOP (32.4 mg, 62.4 μ mol), DIEA (21.4 mg, 166.4 μ mol) and (S) ethyl lactate (19.6 mg, 166.4 μ mol). The reaction mixture was stirred at room temperature for one hour. The crude reaction mixture was purified by RP HPLC on C₁₈ column using H₂O/acetonitrile (5-95%) to provide 13.5 mg (81%) of the pure product as a mixture of diastereomers at phosphorus (~ 4:1). ¹H NMR (300 MHz, CDCl₃) δ 0.0 (s, 9H), 1.02 (t, 2H, *J*= 8.7 Hz), 1.23 (t, 3H, *J*= 9.3 Hz), 1.35 (d, 2.4H, *J*= 6.6 Hz), 1.42 – 1.53 (m, 4.6H), 1.67 – 1.86 (m, 4H), 3.14 (s, 3H), 4.03 – 4.27 (m, 4H), 4.71 (br s, 3H), 4.98 (m, 0.8H), 5.10 (m, 0.2H), 6.57 (d, 2H, *J*= 7.5 Hz), 7.00 (m, 1H), 7.16 (m, 3H), 7.30 (m, 2H), 7.63 (d, 2H, *J*= 7.5 Hz), 8.43 (s, 1H); ³¹P (121.4 MHz, DMSO-*d*₆) δ 30.5, 29.2; MS (*m/z*) 795.2 [M]⁺.

Example 348

By way of example and not limitation, embodiments of the invention are named below in tabular format (Table 100). These embodiments are of the general formula "MBF":

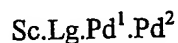


MBF

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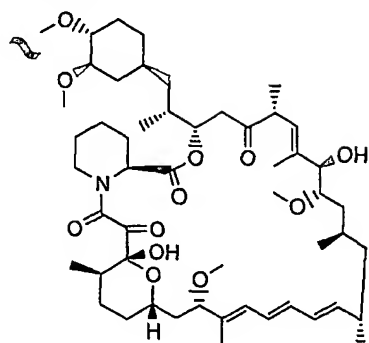
Each embodiment of MBF is depicted as a substituted nucleus (Sc). Sc is described in formula 1-151 herein, wherein A⁰ is the point of covalent attachment of Sc to Lg, as well as in Tables 1.1 to 1.5 below. For those
 10 embodiments described in Table 100, Sc is a nucleus designated by a number and each substituent is designated in order by letter or number. Tables 1.1 to 1.5 are a schedule of nuclei used in forming the embodiments of Table 100. Each nucleus (Sc) is given a number designation from Tables 1.1 to 1.5, and this designation appears first in each embodiment name. Similarly, Tables 10.1 to
 15 10.19 and 20.1 to 20.36 list the selected linking groups (Lg) and prodrug (Pd¹ and Pd²) substituents, again by letter or number designation, respectively. Accordingly, a compound of the formula MBF includes compounds having Sc groups based on formula 1-151 herein as well as compounds according to Table 100 below. In all cases, compounds of the formula MBF have groups Lg, Pd¹
 20 and Pd² setforth in the Tables below.

Accordingly, each named embodiment of Table 100 is depicted by a number designating the nucleus from Table 1.1-1.5, followed by a letter designating the linking group (Lg) from Table 10.1-10.19, and two numbers designating the two prodrug groups (Pd¹ and Pd²) from Table 20.1-20.36. In
 25 graphical tabular form, each embodiment of Table 100 appears as a name having the syntax:

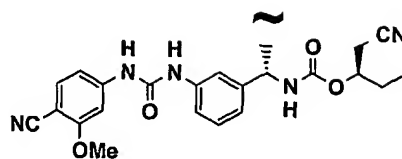


Each Sc group is shown having a tilda ("~"). The tilda is the point of covalent attachment of Sc to Lg. Q¹ and Q² of the linking groups (Lg), it should be understood, do not represent groups or atoms but are simply connectivity designations. Q¹ is the site of the covalent bond to the nucleus (Sc) and Q² is the site of the covalent bond to the phosphorous atom of formula MBF. Each prodrug group (Pd¹ and Pd²) are covalently bonded to the phosphorous atom of MBF at the tilda symbol ("~"). Some embodiments of Tables 10.1-10.19 and 20.1-20.36 may be designated as a combination of letters and numbers (Table 10.1-10.19) or number and letter (Table 20.1-20.36). For example there are Table 10 entries for BJ1 and BJ2. In any event, entries of Table 10.1-10.19 always begin with a letter and those of Table 20.1-20.36 always begin with a number. When a nucleus (Sc) is shown enclosed within square brackets ("[]") and a covalent bond extends outside the brackets, the point of covalent attachment of Sc to Lg may be at any substitutable site on SC. Selection of the point of attachment is described herein. By way of example and not limitation, the point of attachment is selected from those depicted in the schemes and examples.

20

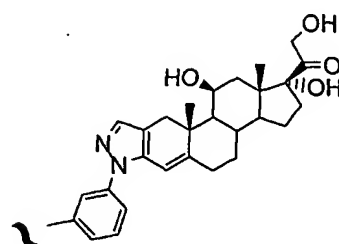
Table 1.1

1

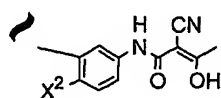


3

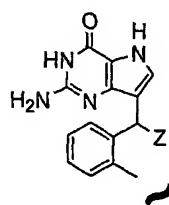
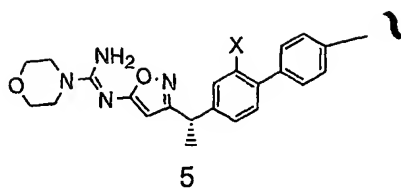
5



2

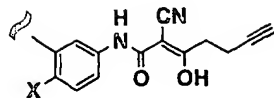


4

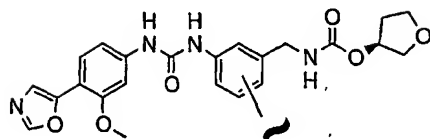
Table 1.2

Z = H, CH₂OH

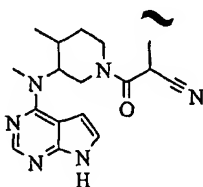
5

Table 1.3

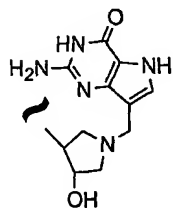
10



9

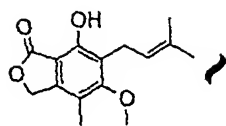


8

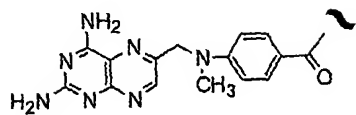


7

5

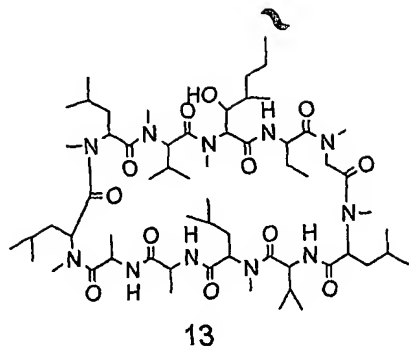
Table 1.4

11



12

5

Table 1.5

5

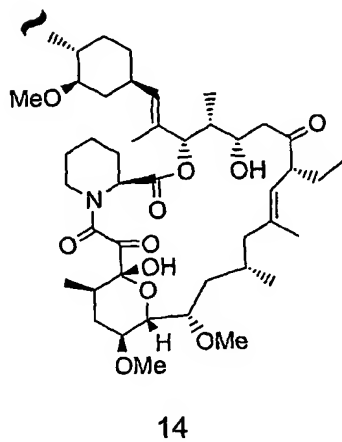
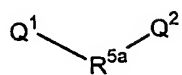
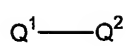
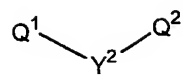


Table 10.1

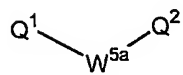
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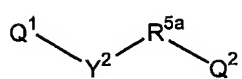
B



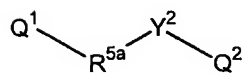
C



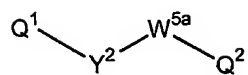
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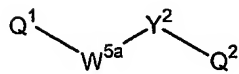
E



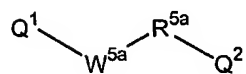
F



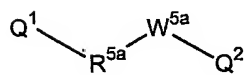
G



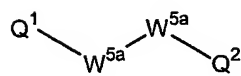
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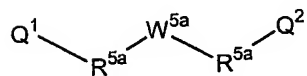
I



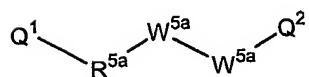
J



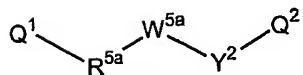
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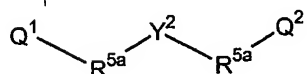
L



M



N



O

Table 10.2

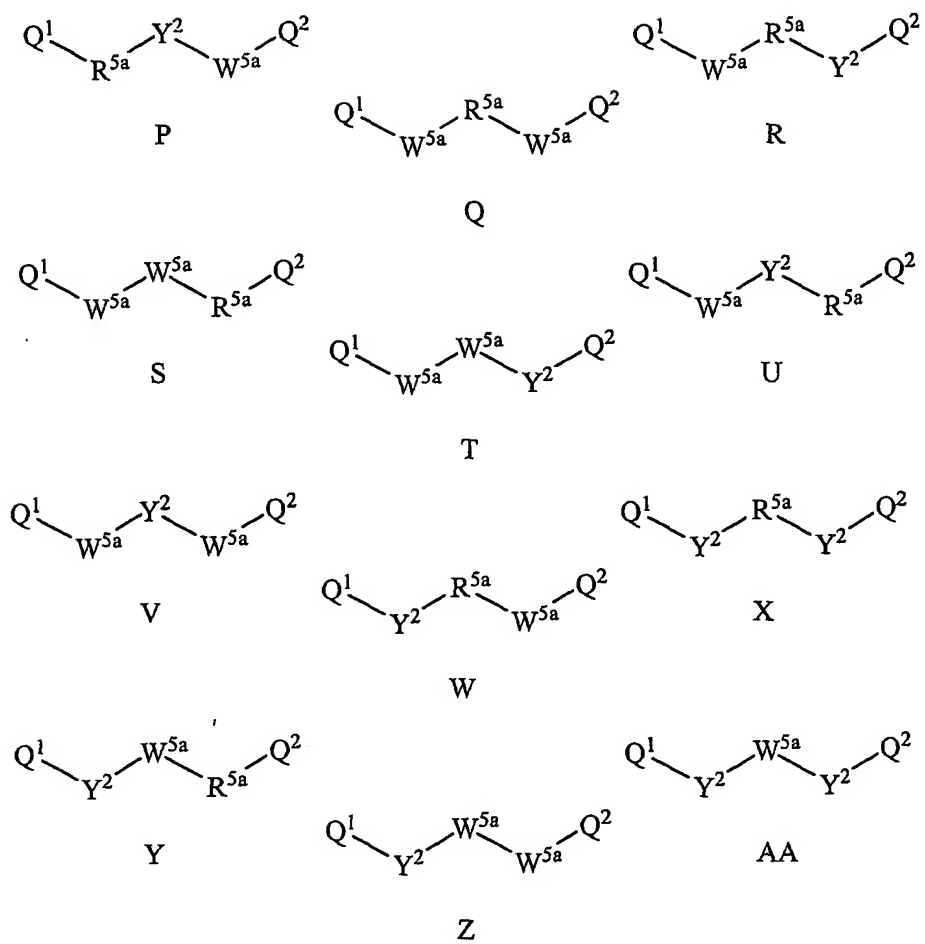


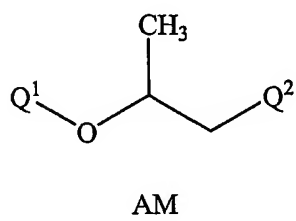
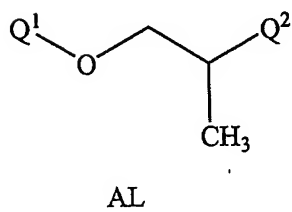
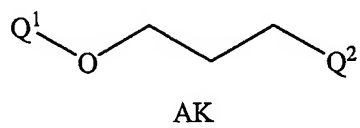
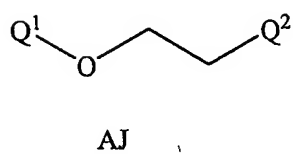
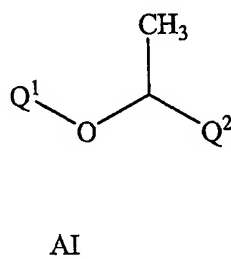
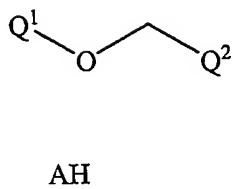
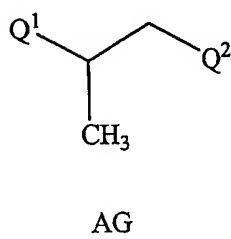
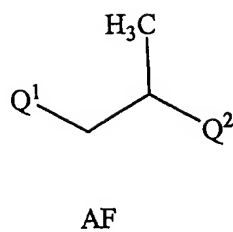
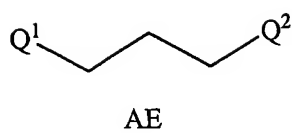
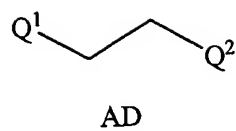
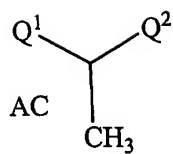
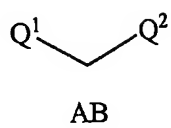
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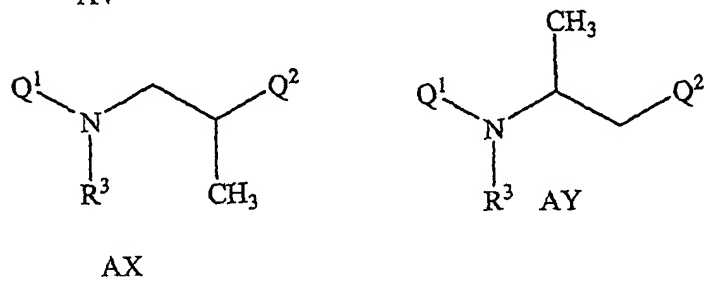
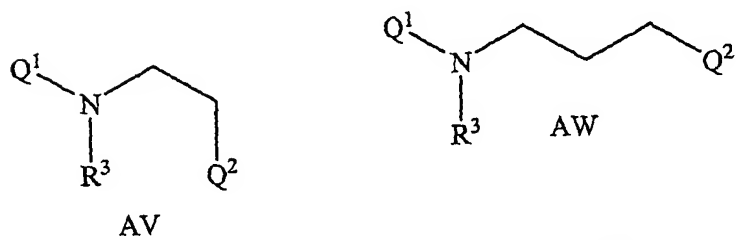
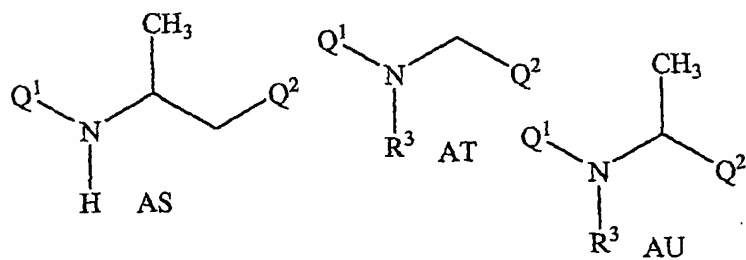
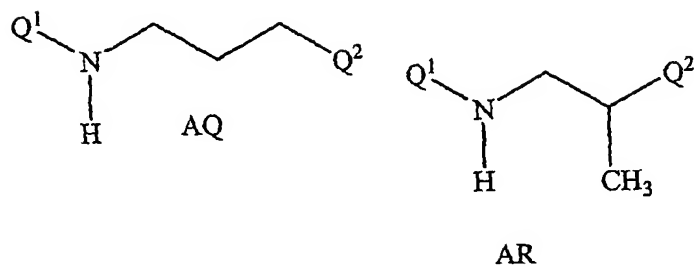
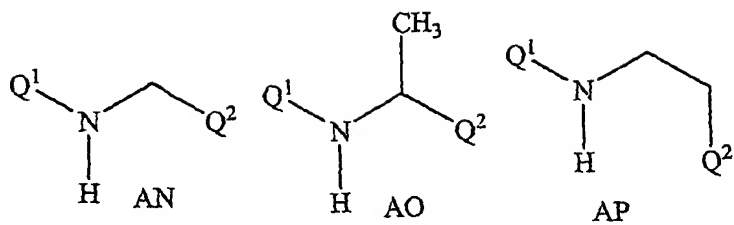
Table 10.4

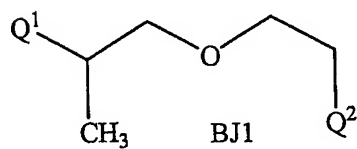
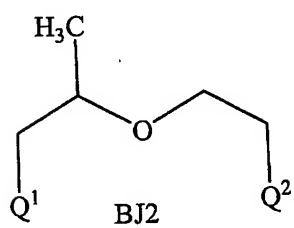
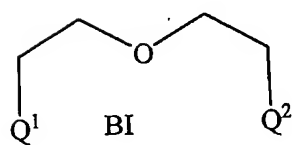
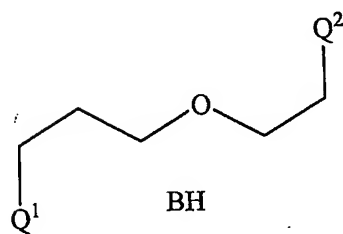
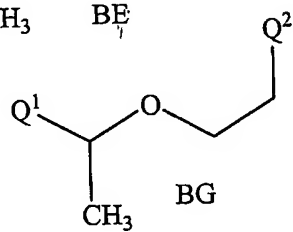
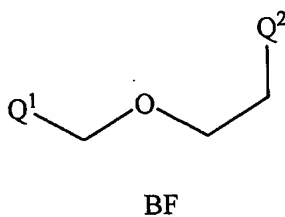
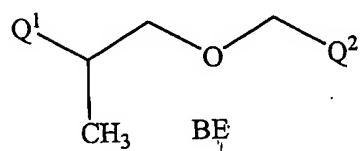
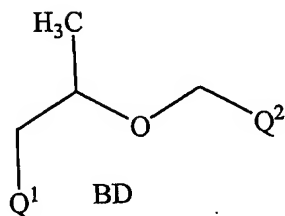
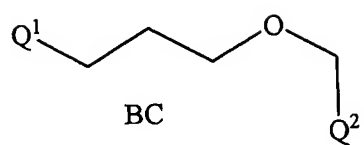
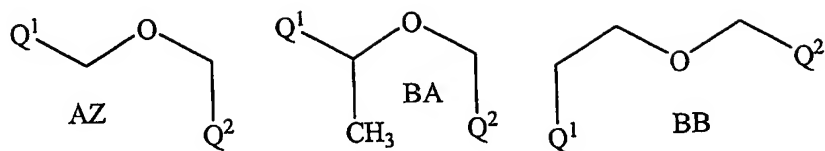
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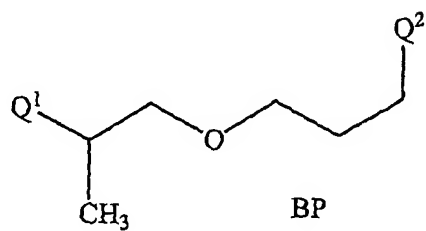
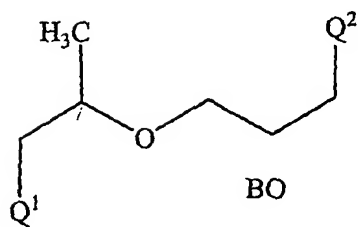
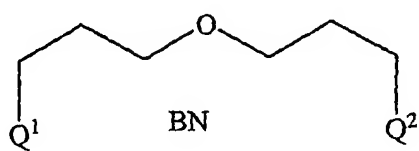
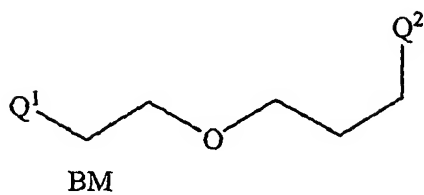
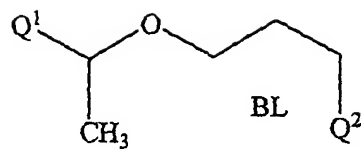
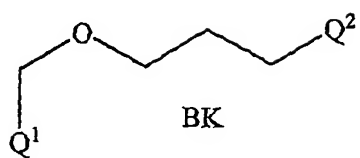
Table 10.6

Table 10.7

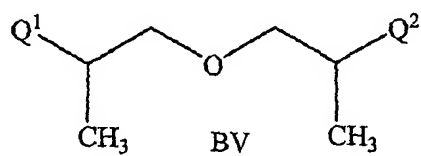
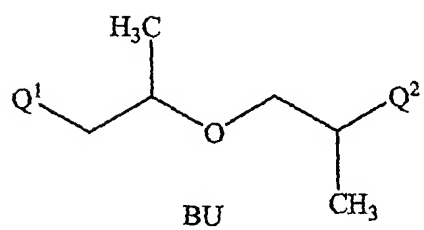
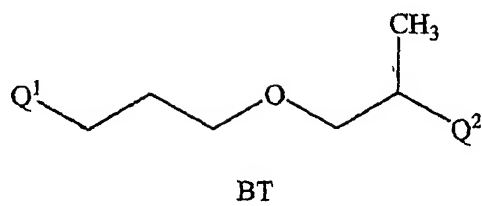
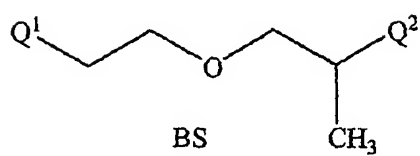
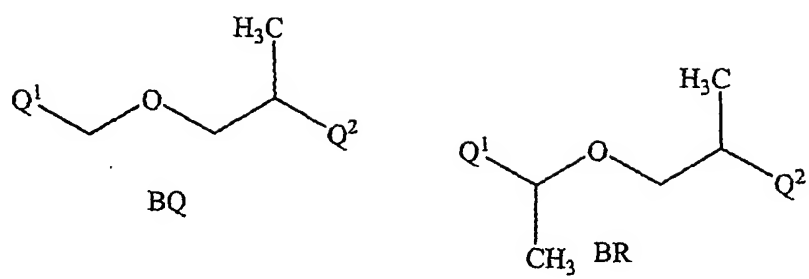


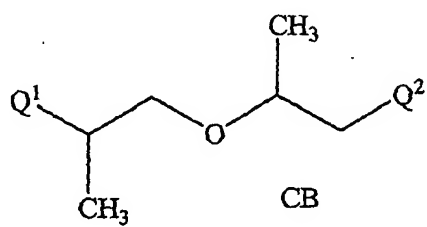
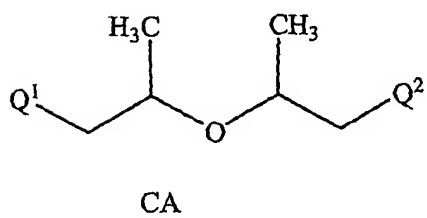
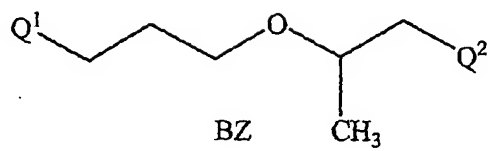
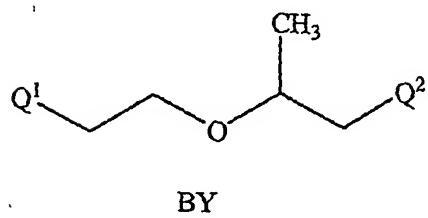
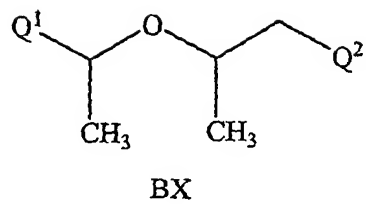
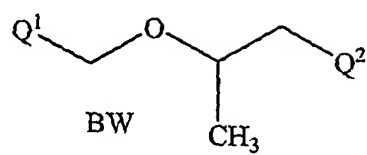
Table 10.8

Table 10.9

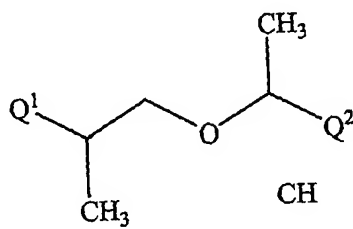
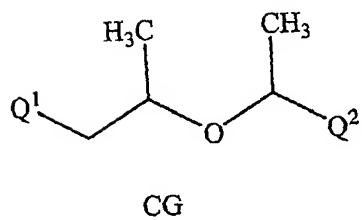
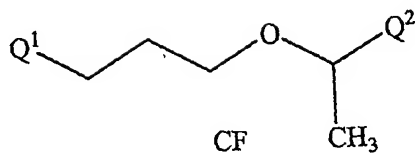
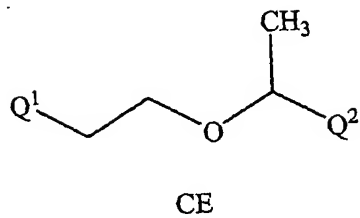
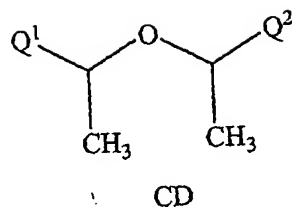
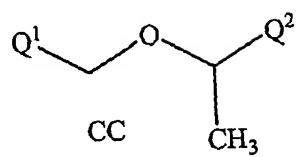


Table 10.10

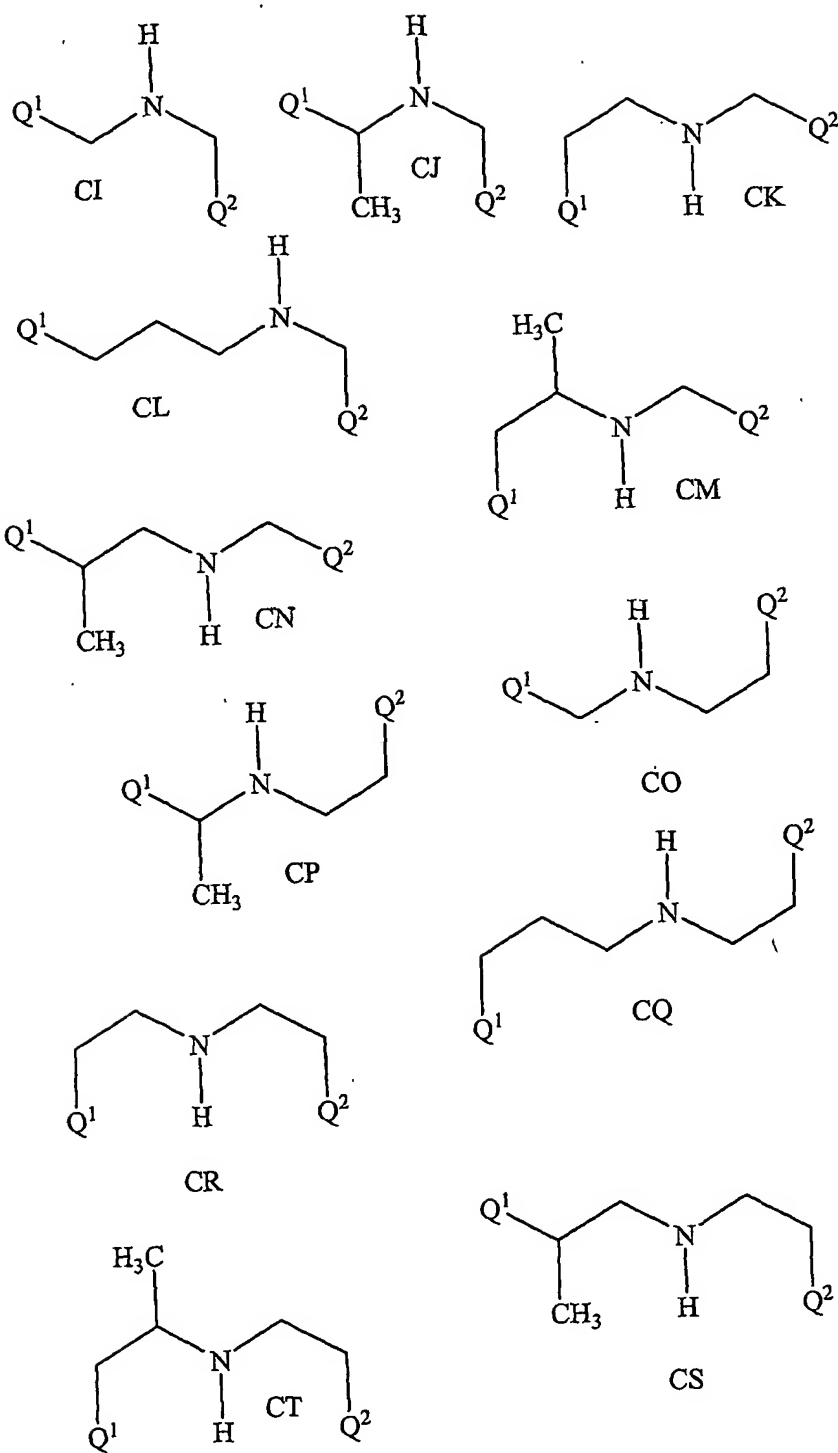


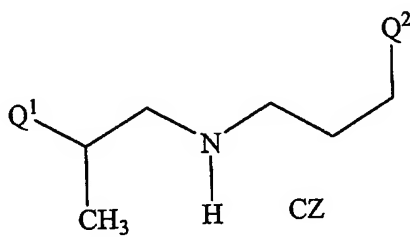
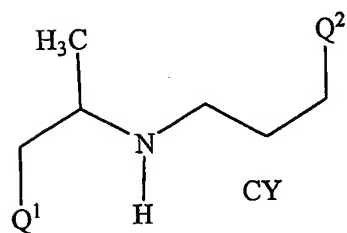
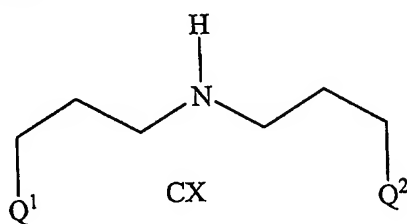
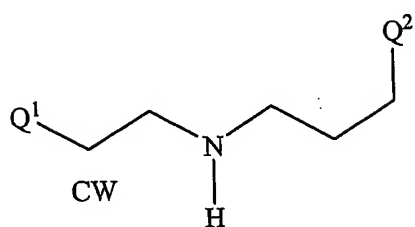
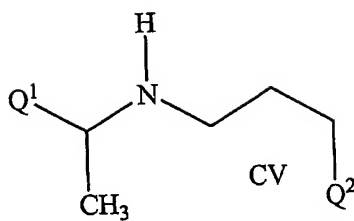
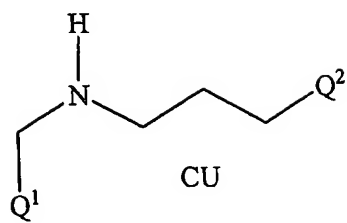
Table 10.11

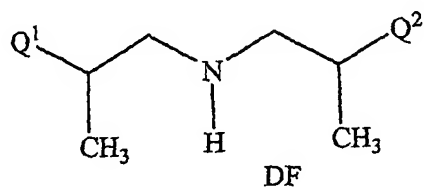
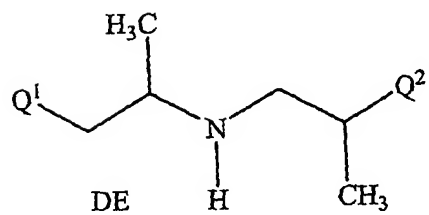
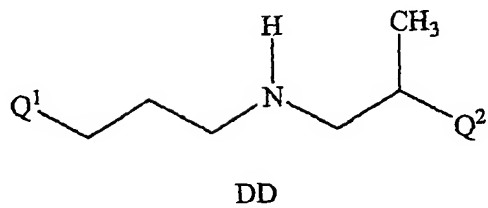
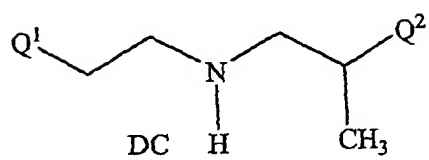
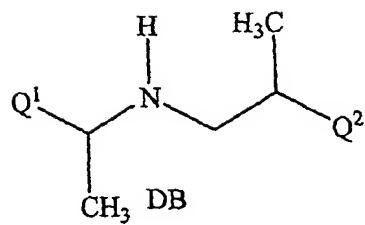
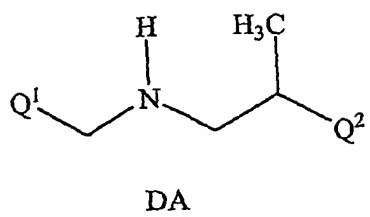
Table 10.12

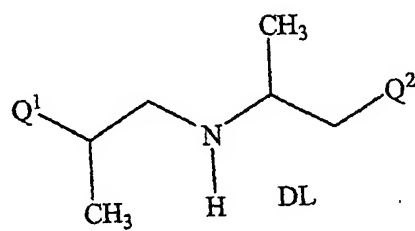
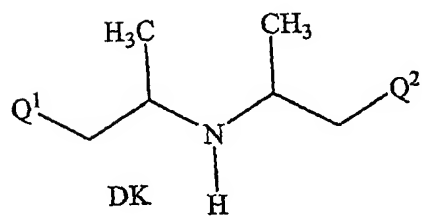
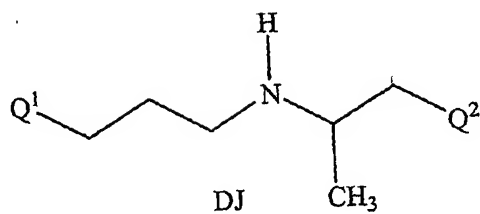
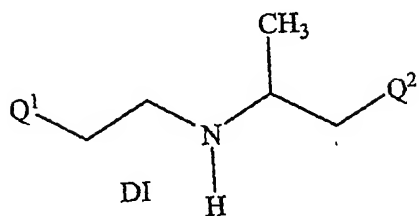
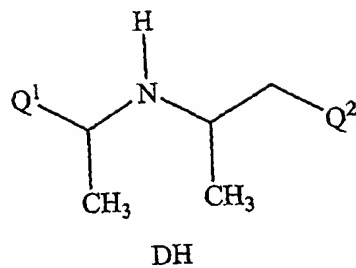
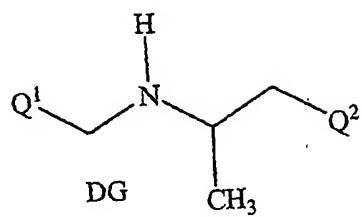
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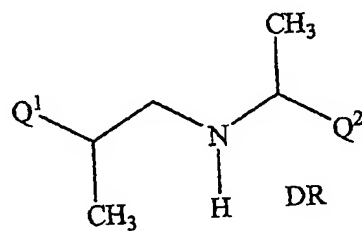
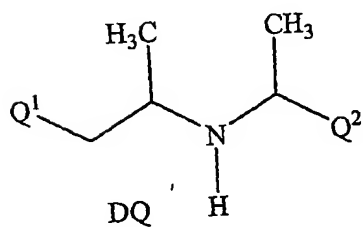
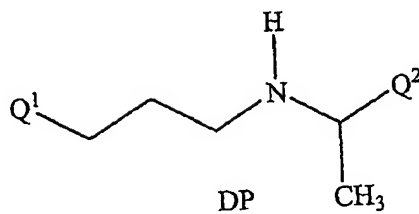
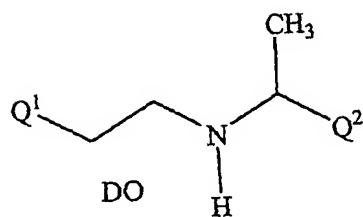
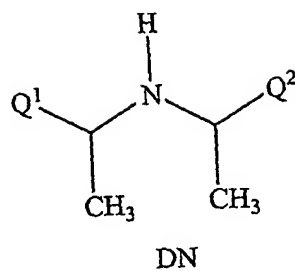
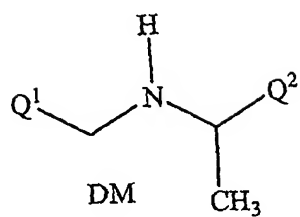
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Table 10.15

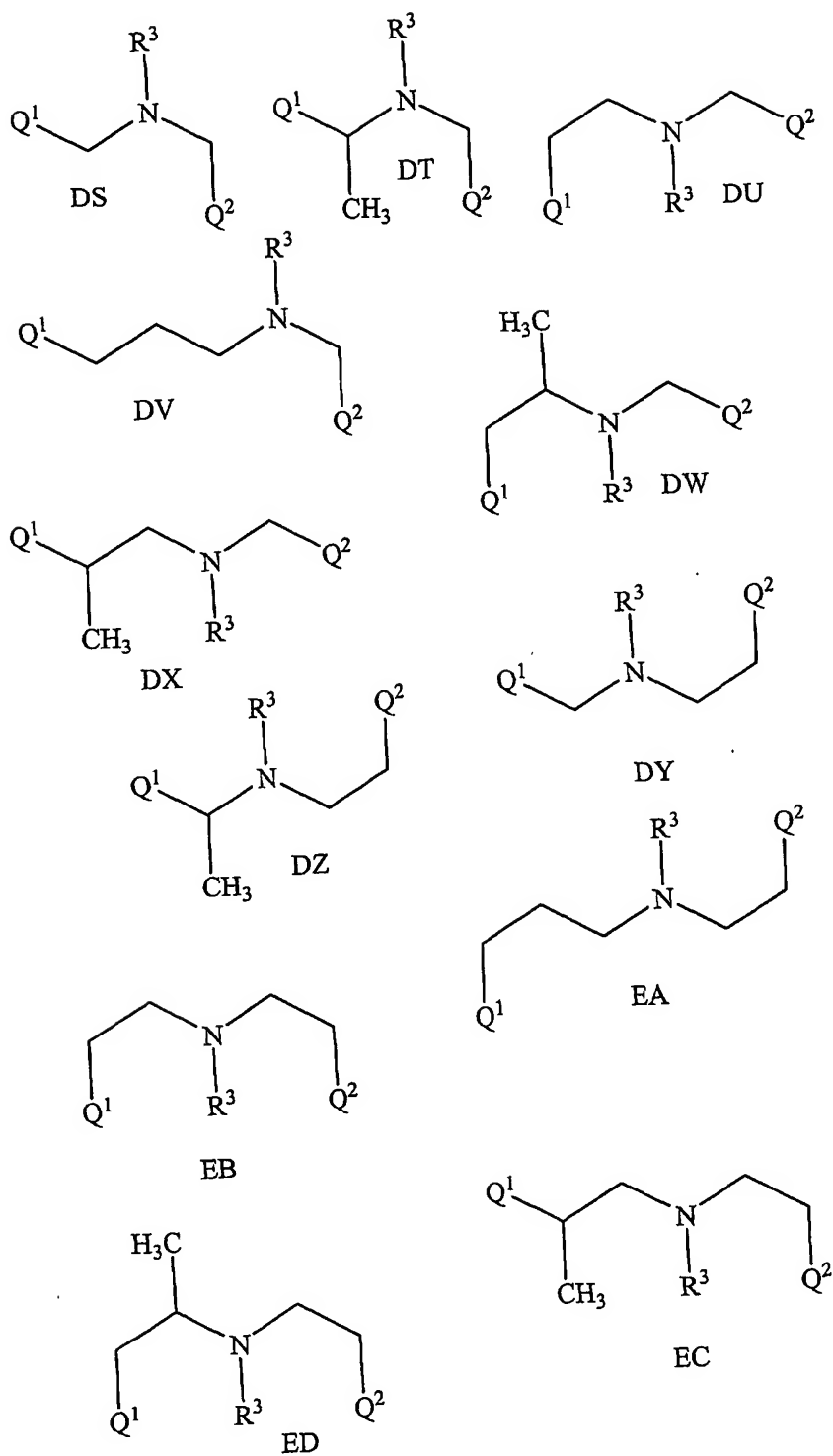


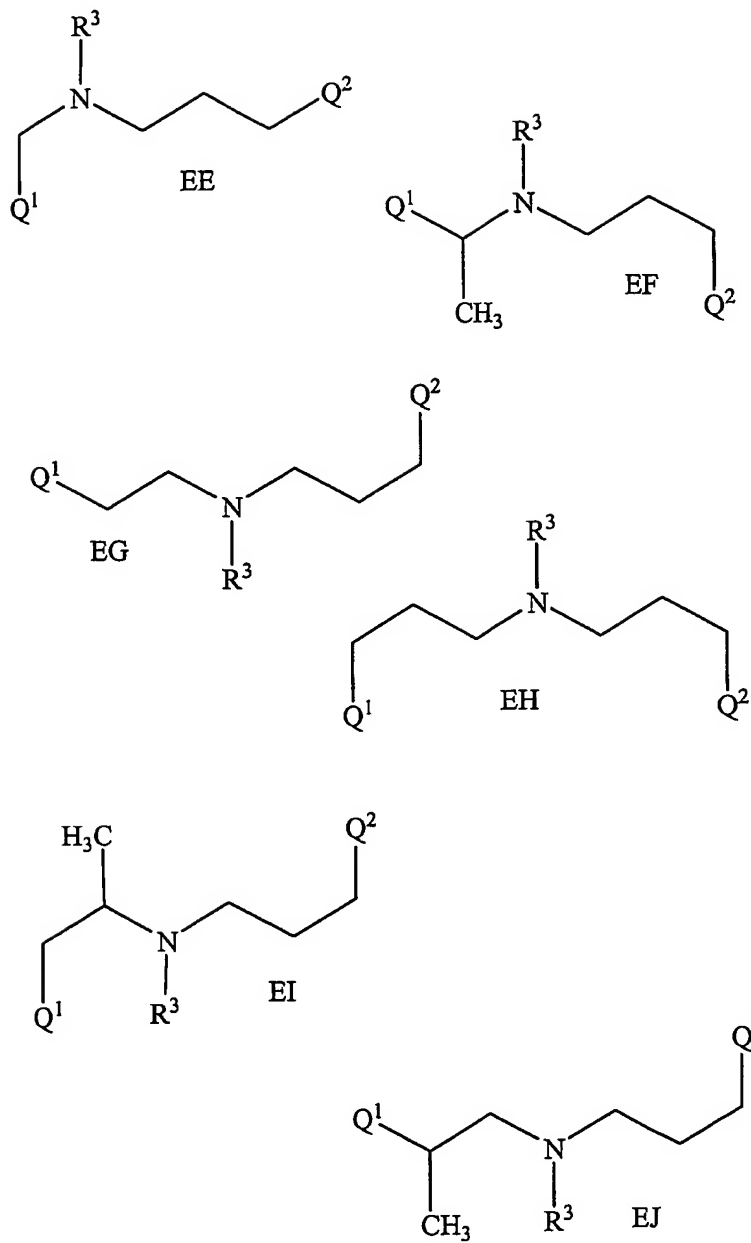
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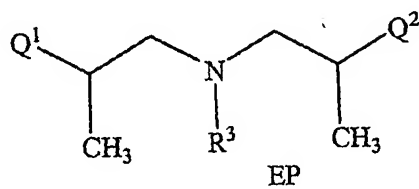
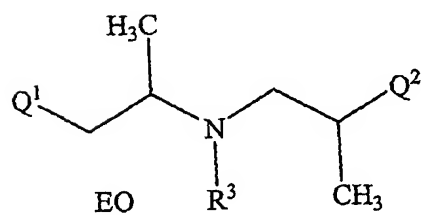
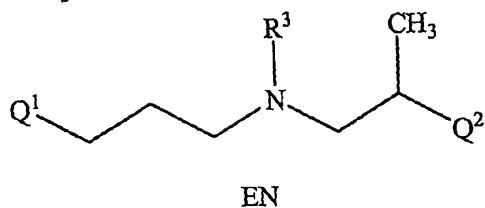
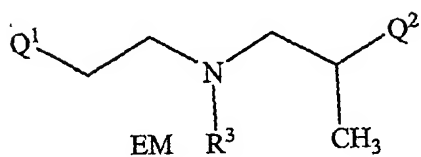
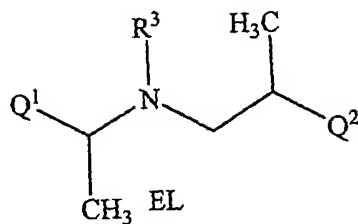
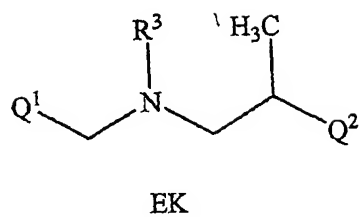
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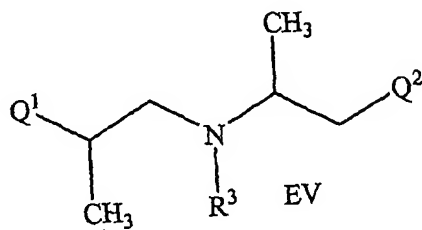
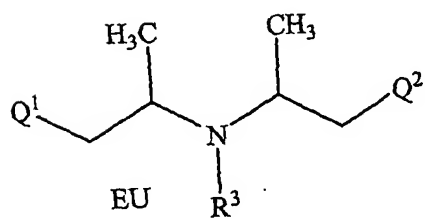
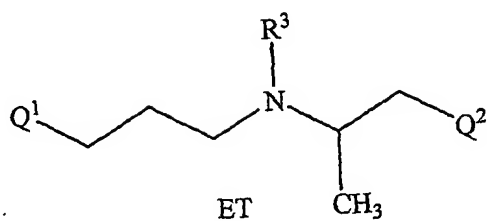
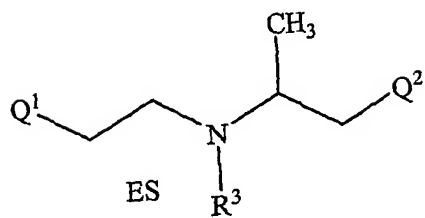
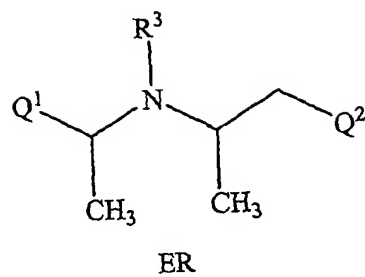
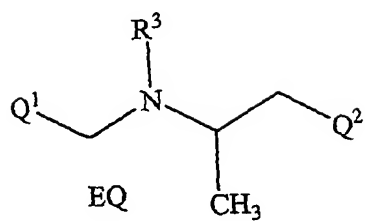
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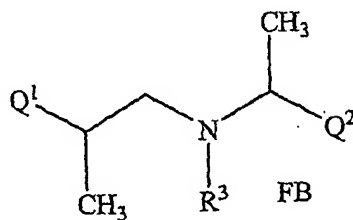
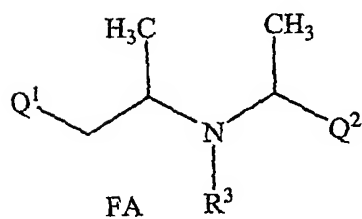
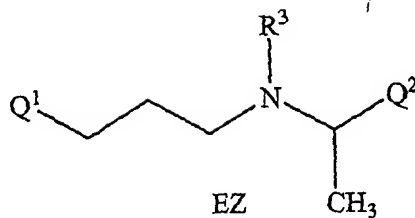
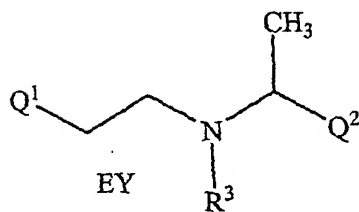
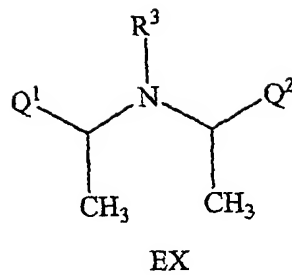
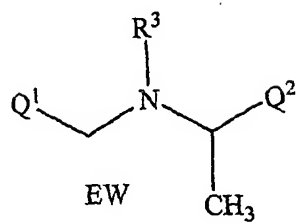
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Table 20.1

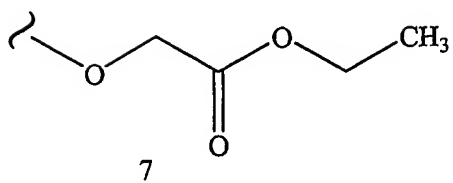
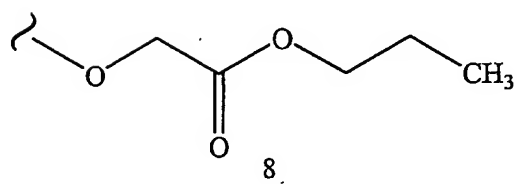
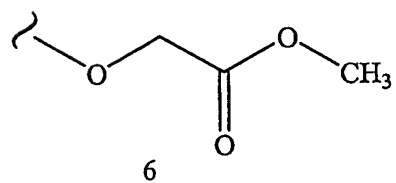
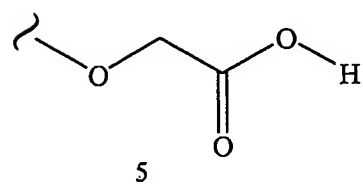
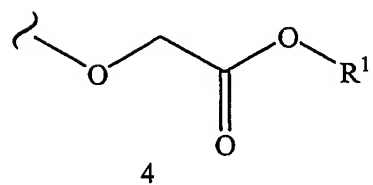
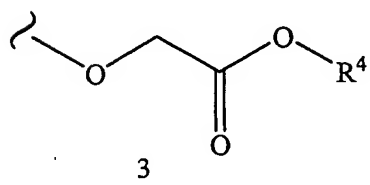
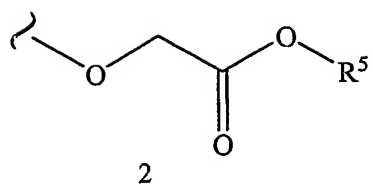
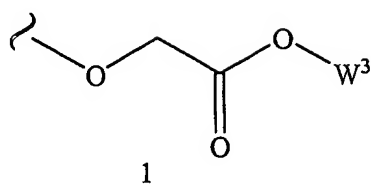


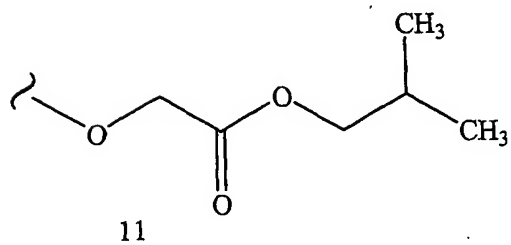
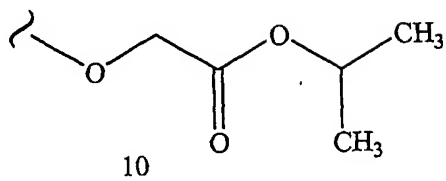
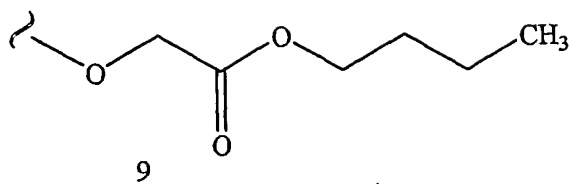
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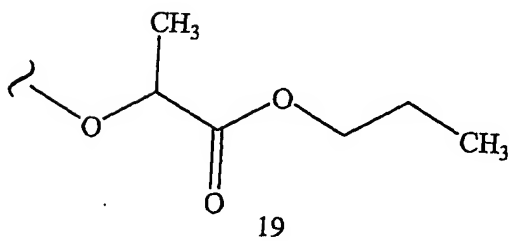
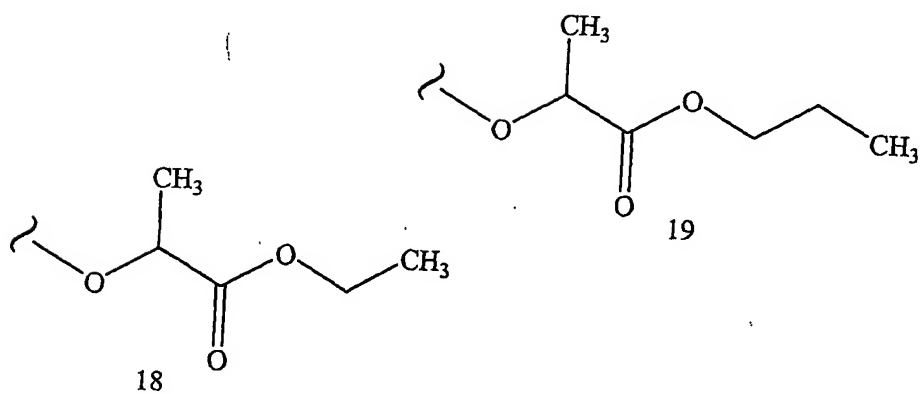
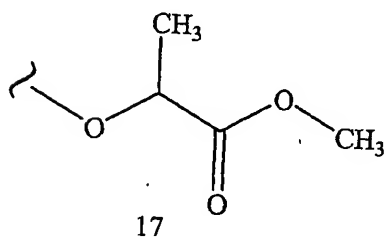
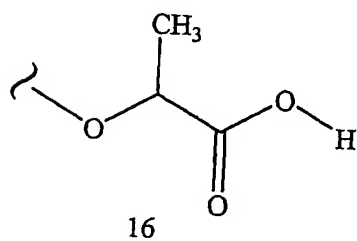
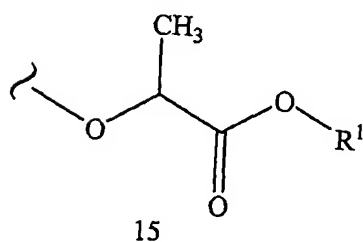
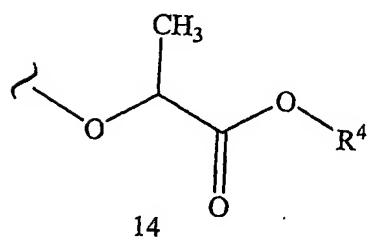
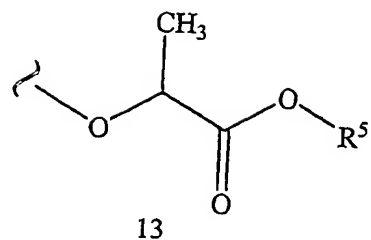
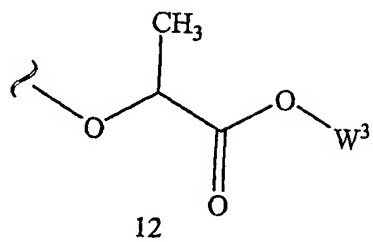
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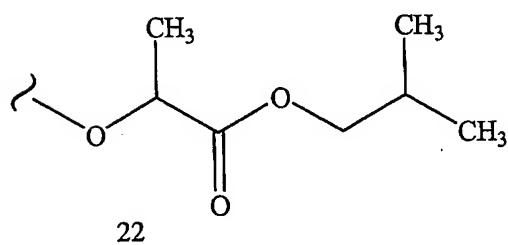
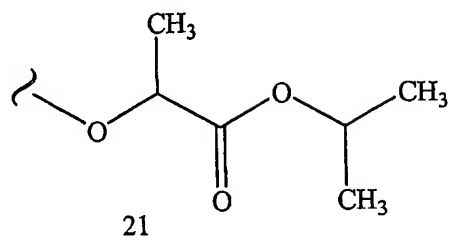
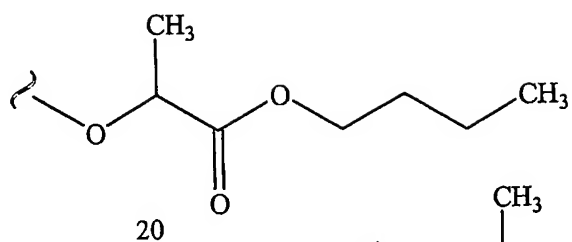
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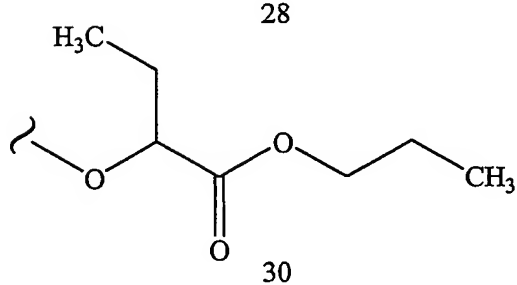
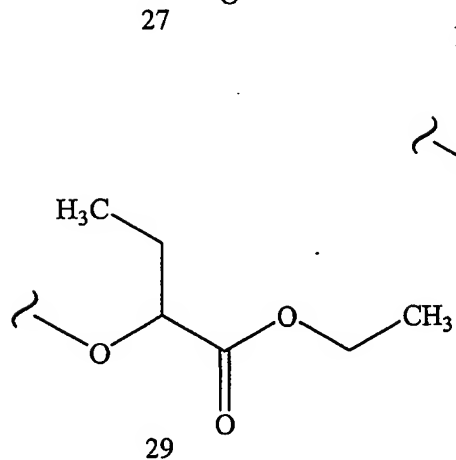
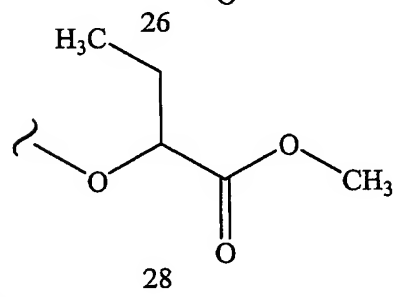
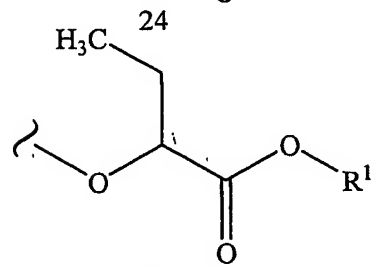
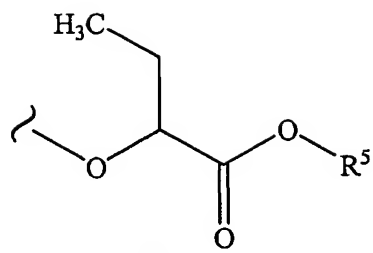
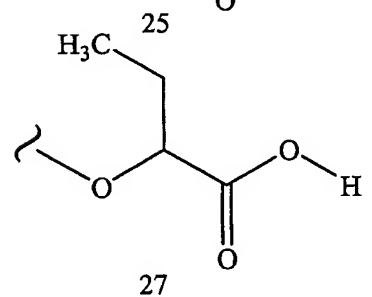
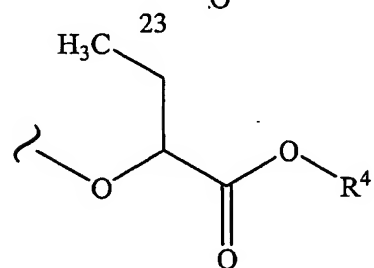
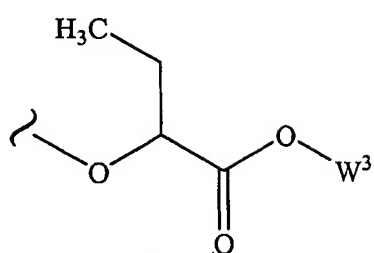
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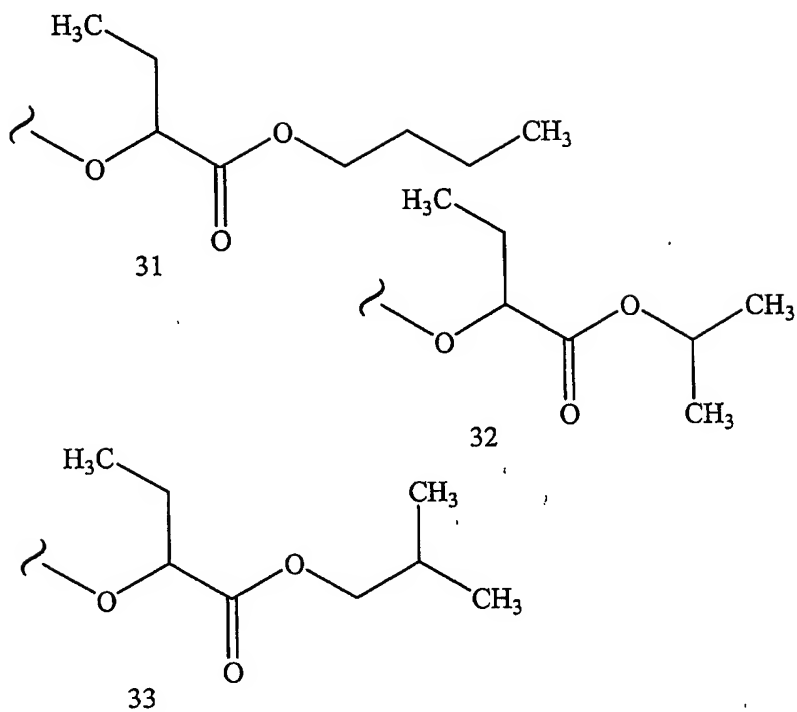
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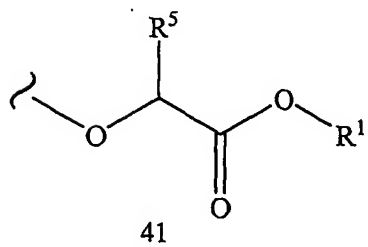
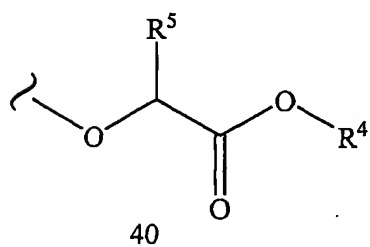
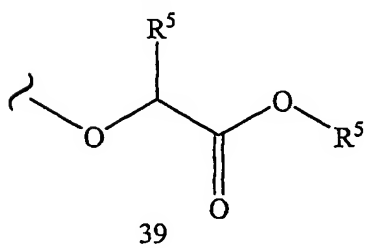
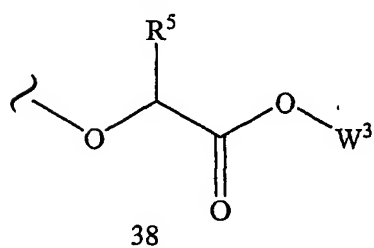
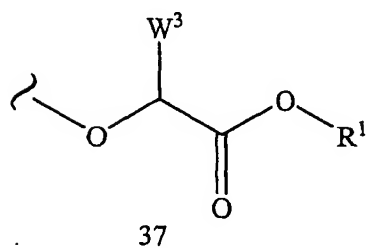
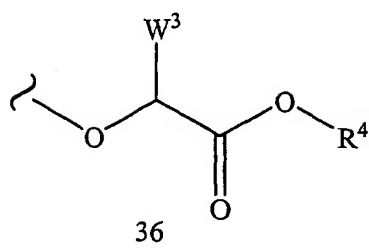
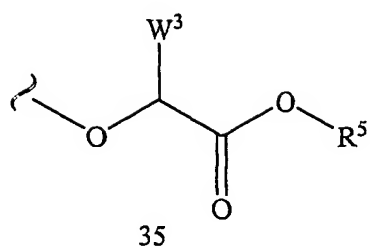
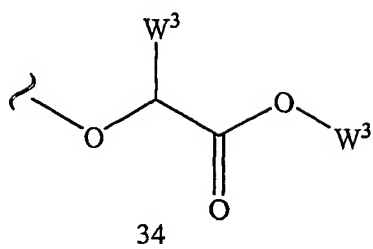
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Table 20.8

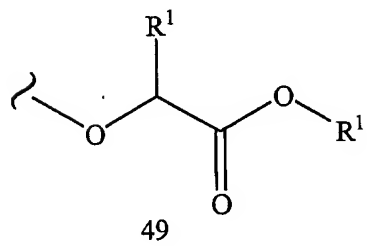
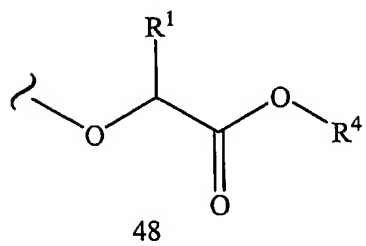
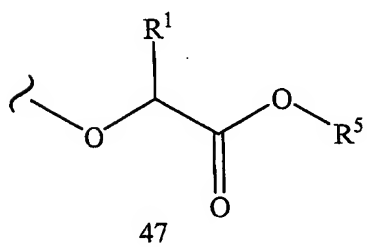
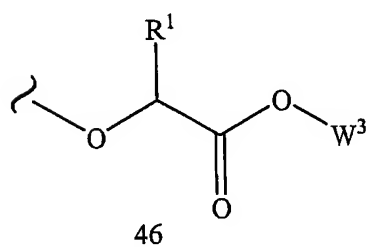
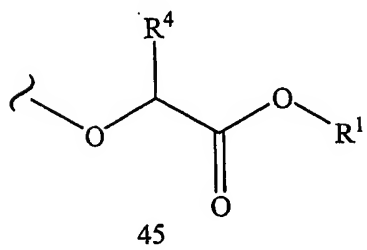
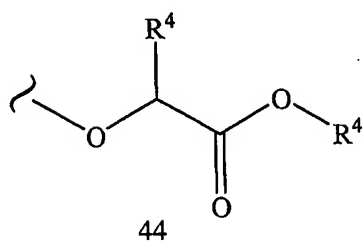
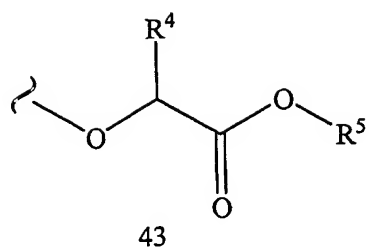
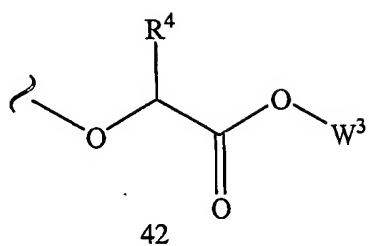


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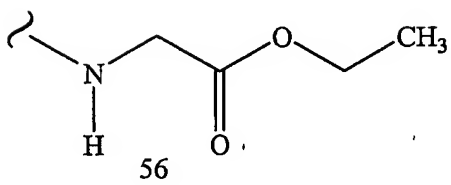
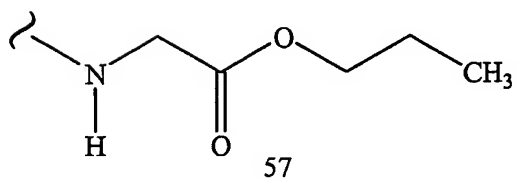
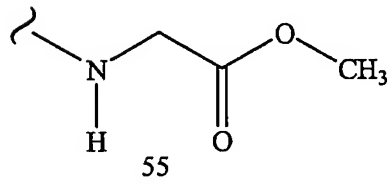
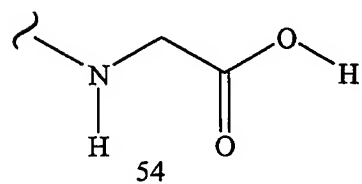
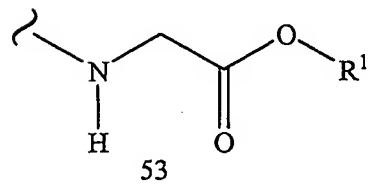
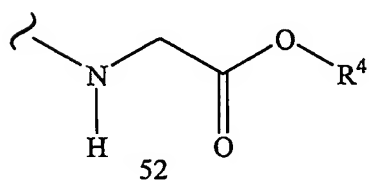
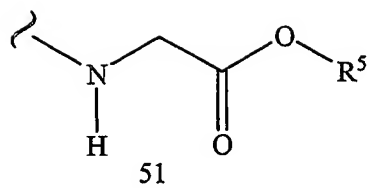
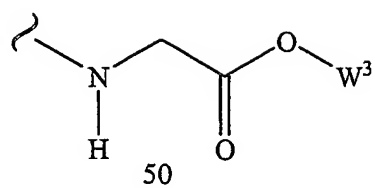


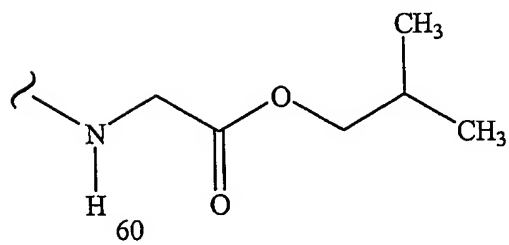
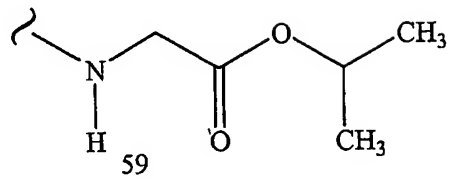
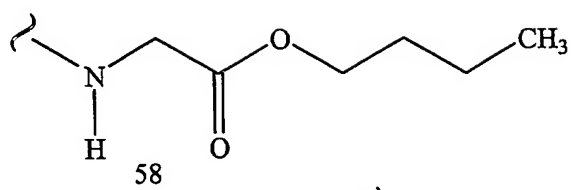
Table 20.10

Table 20.11

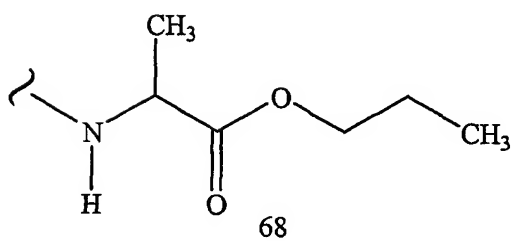
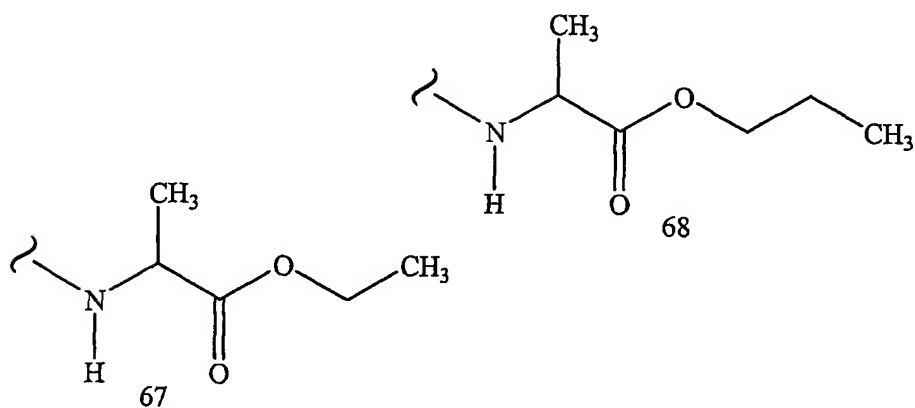
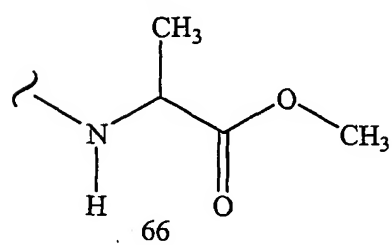
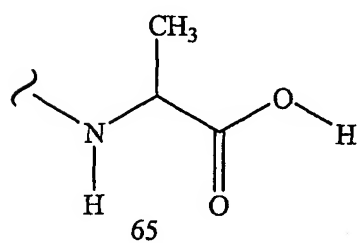
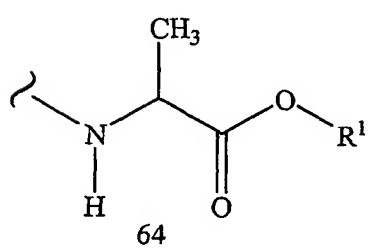
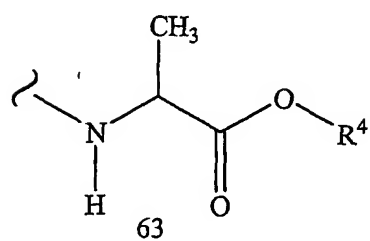
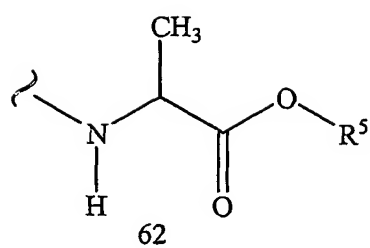
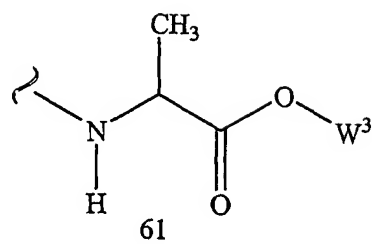


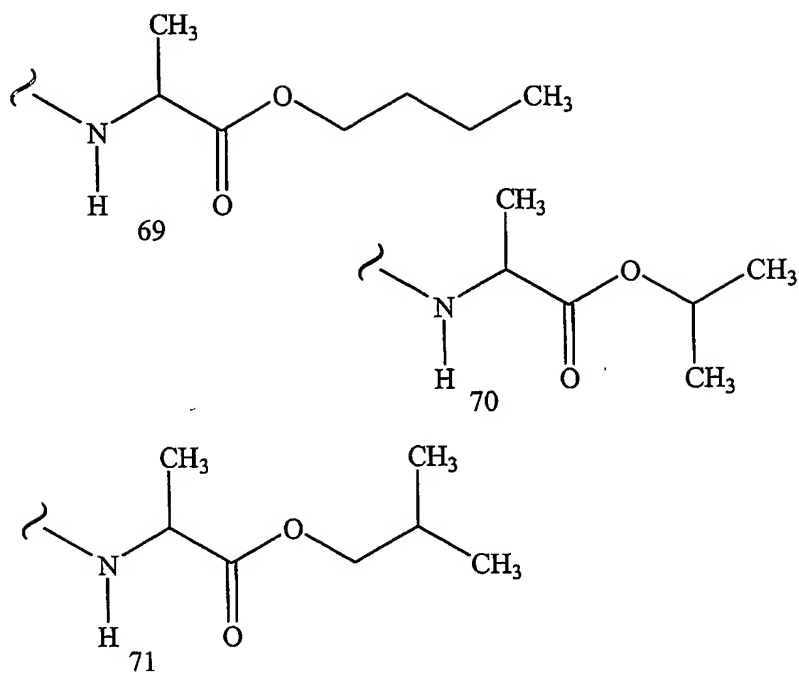
Table 20.12

Table 20.13

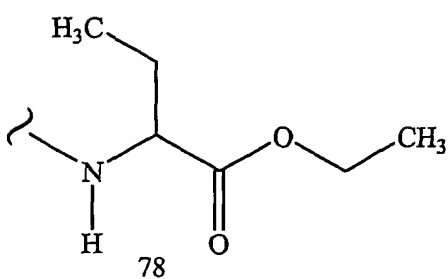
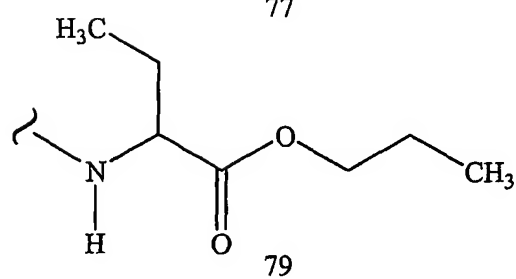
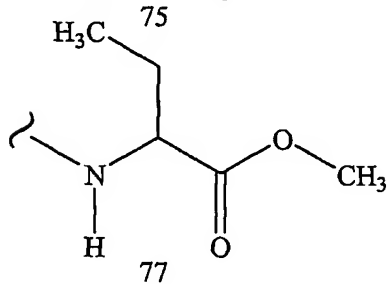
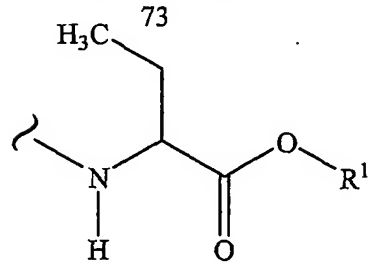
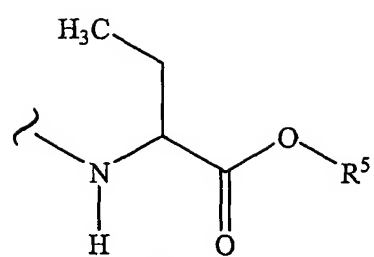
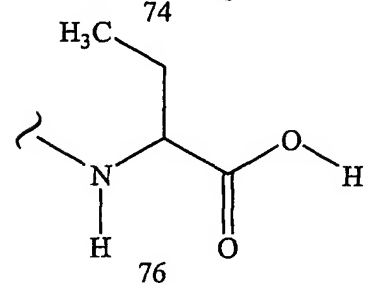
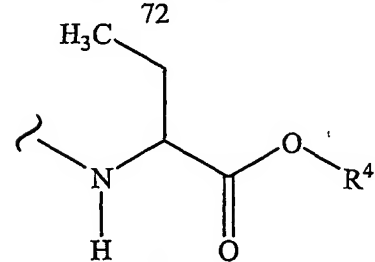
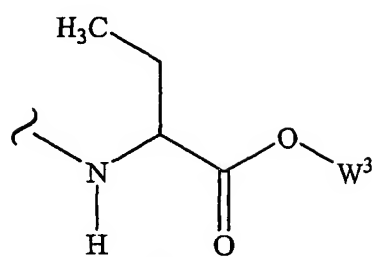


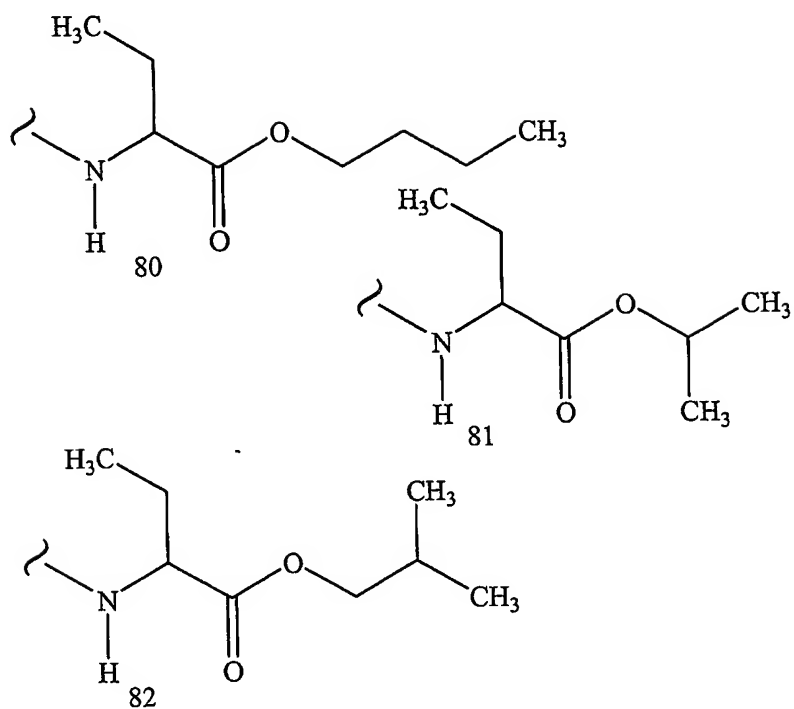
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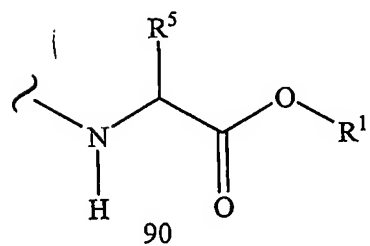
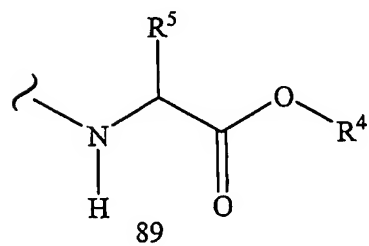
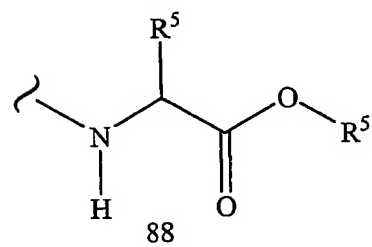
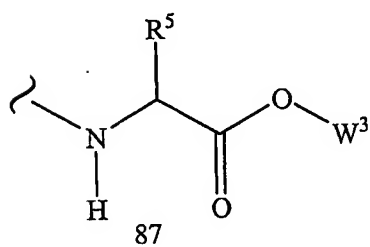
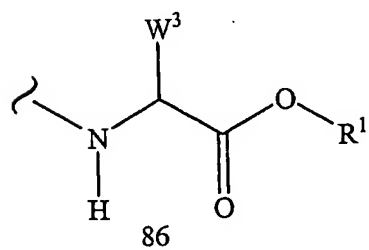
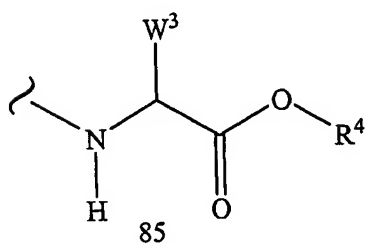
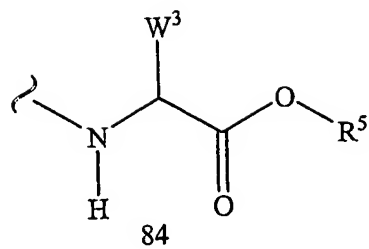
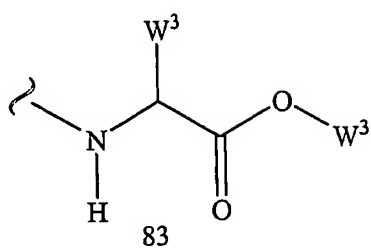
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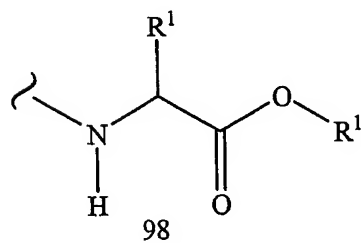
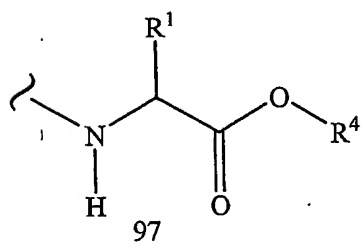
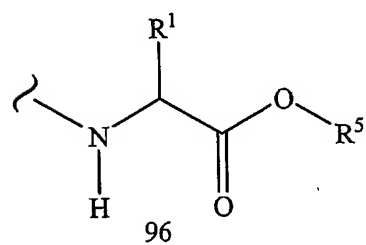
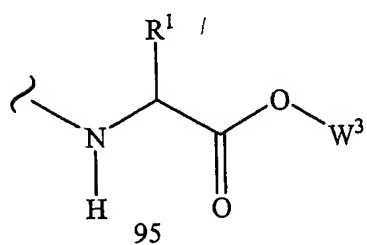
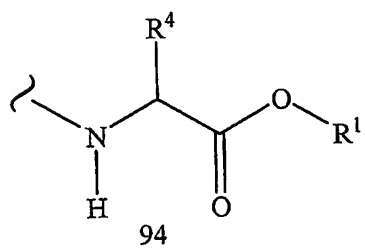
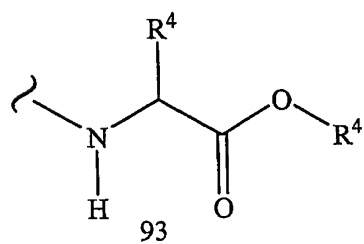
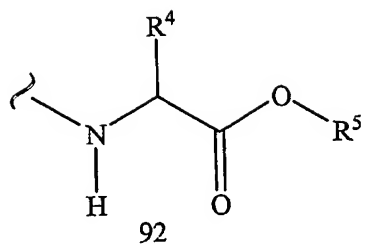
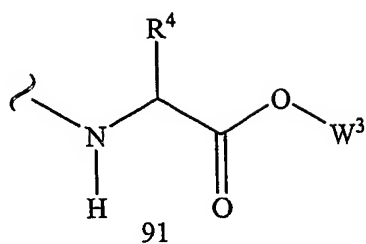
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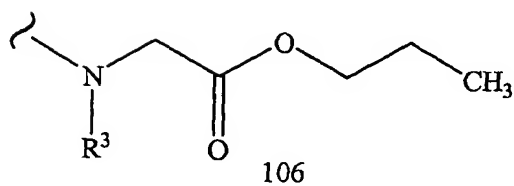
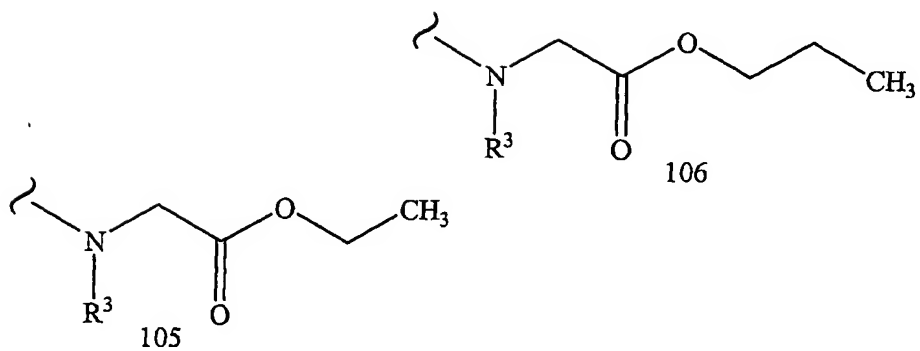
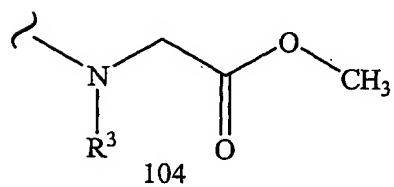
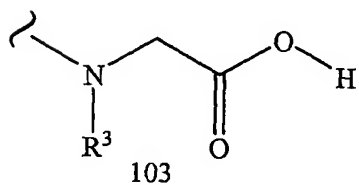
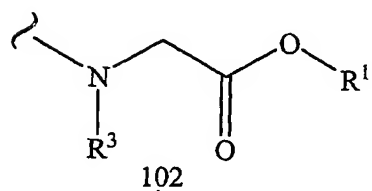
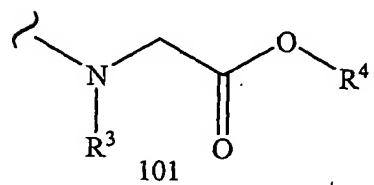
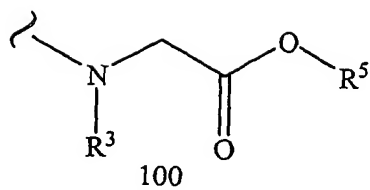
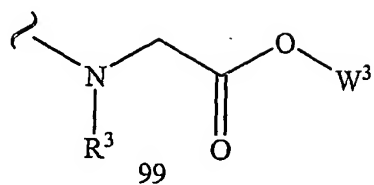
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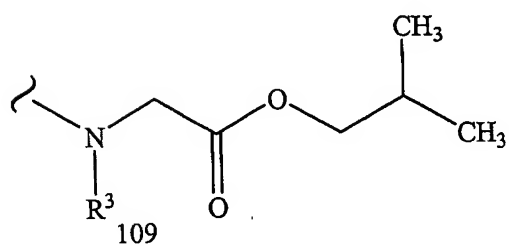
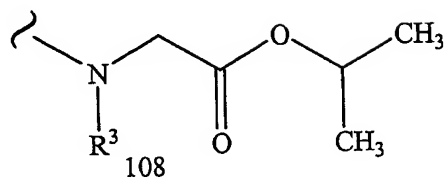
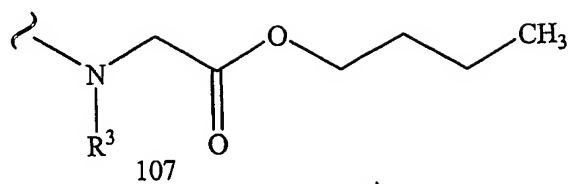
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Table 20.19

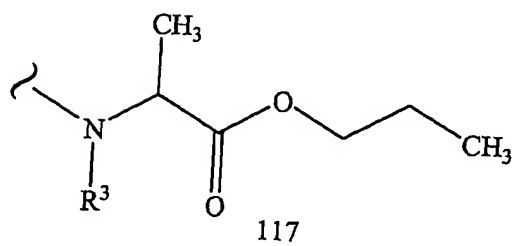
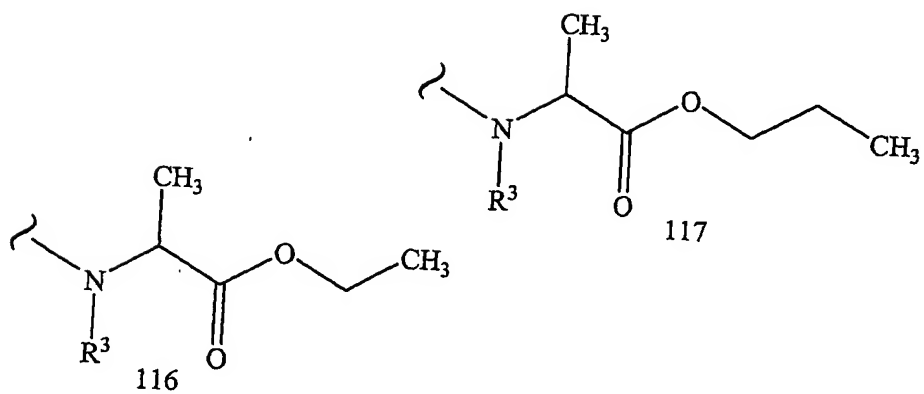
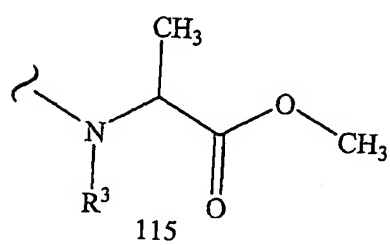
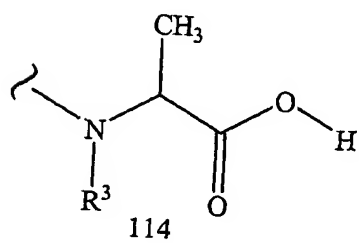
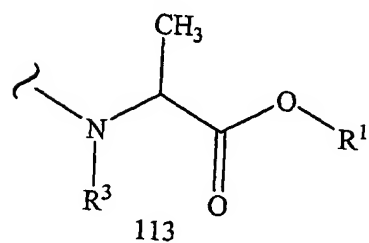
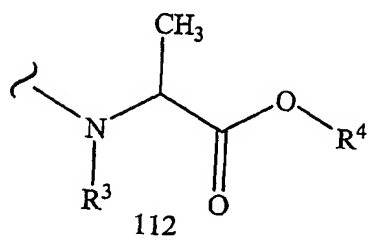
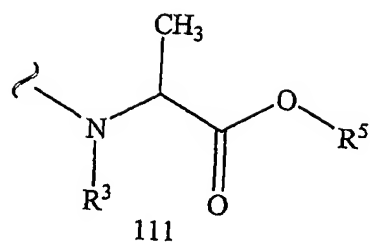
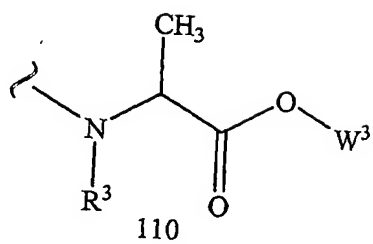


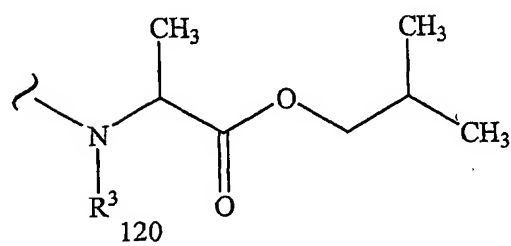
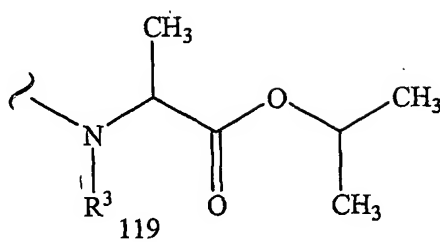
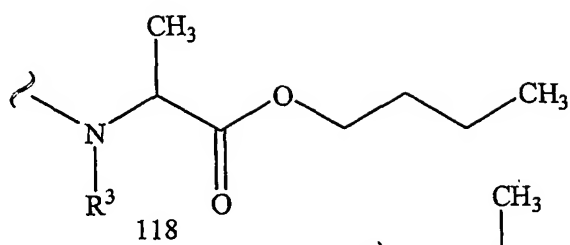
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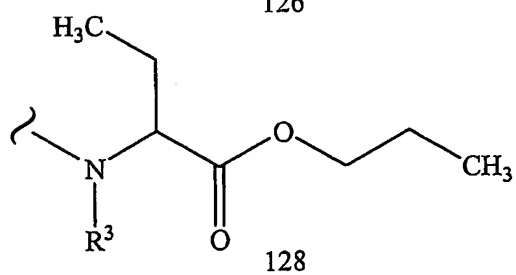
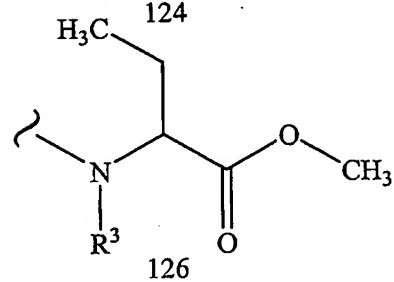
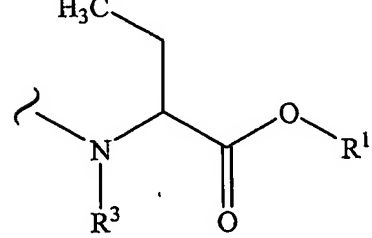
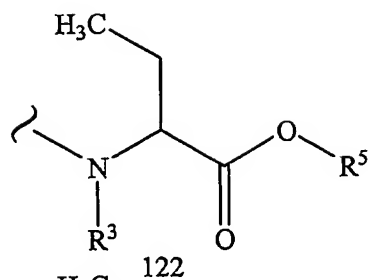
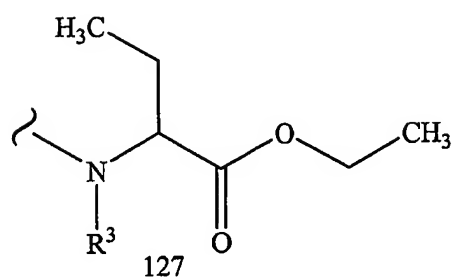
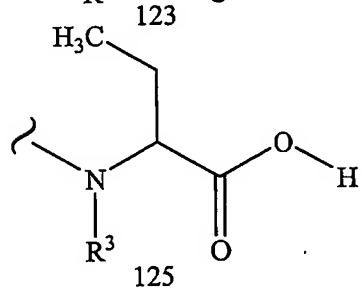
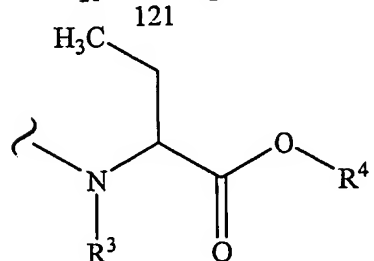
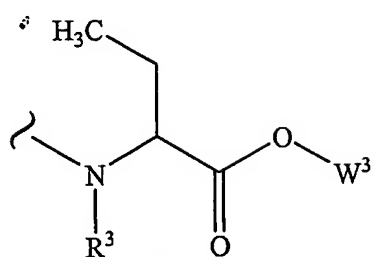
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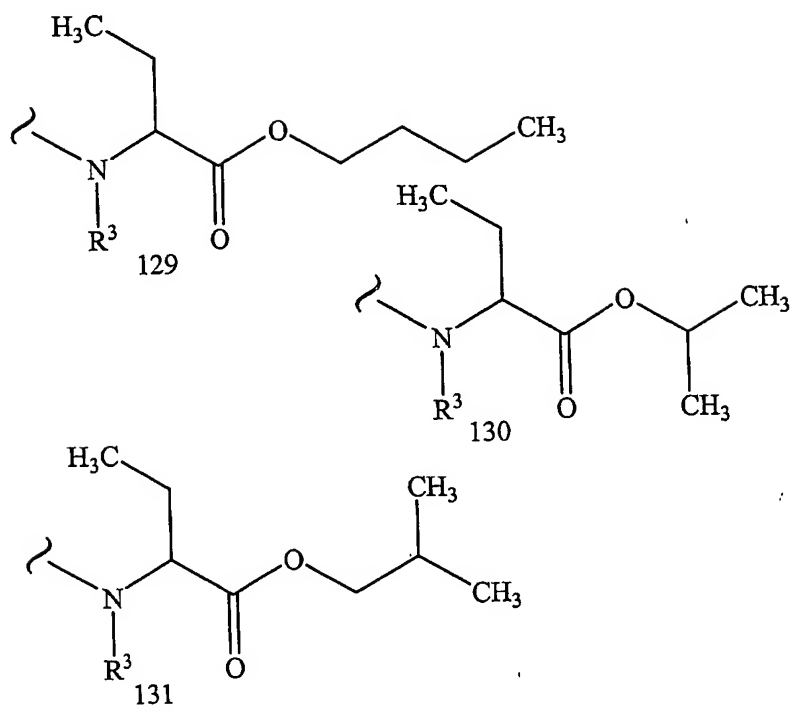
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Table 20.23

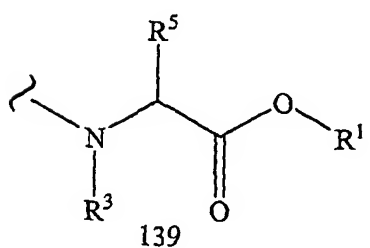
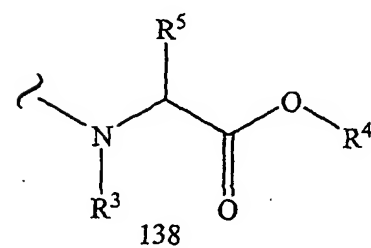
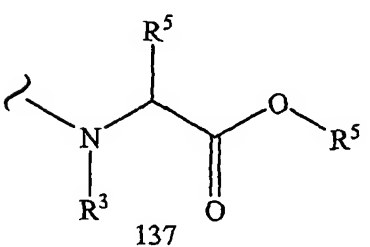
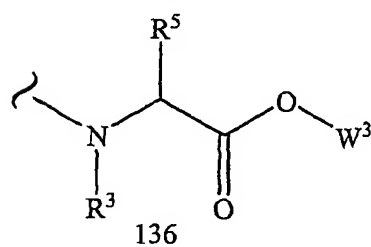
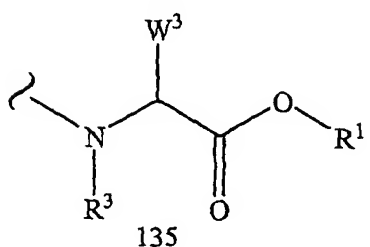
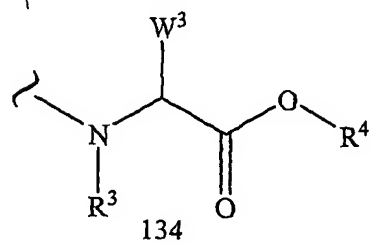
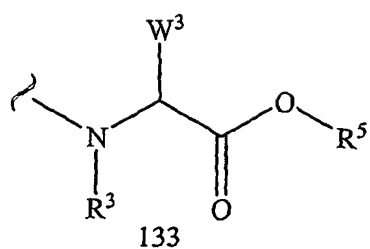
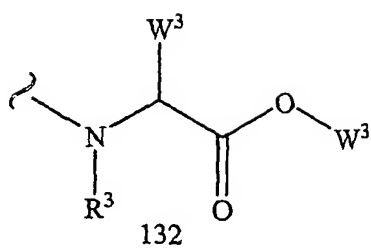


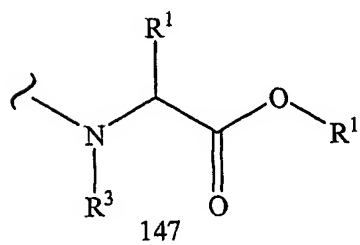
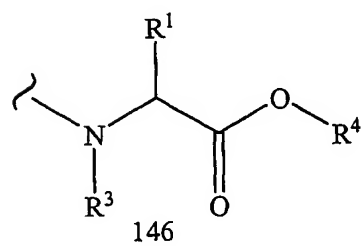
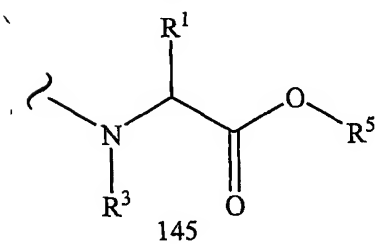
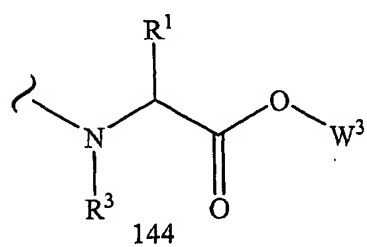
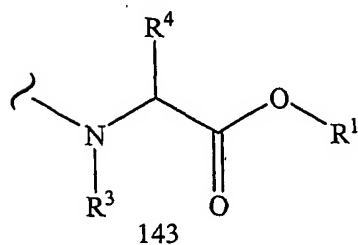
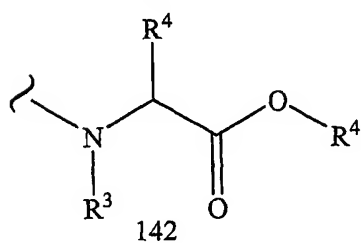
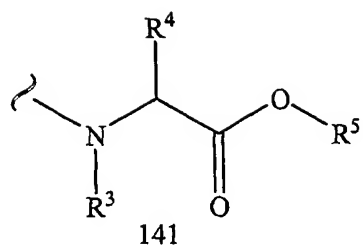
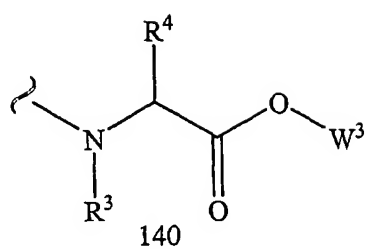
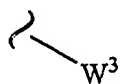
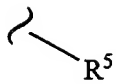
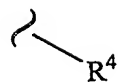
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Table 20.25

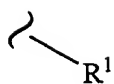
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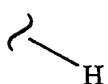
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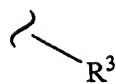
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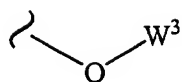
151



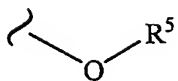
152



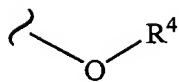
153



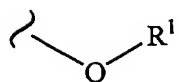
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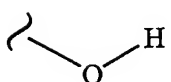
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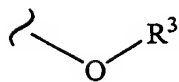
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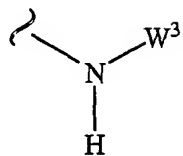
157



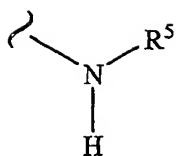
158



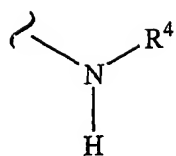
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Table 20.26

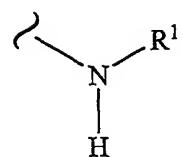
160



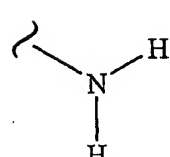
161



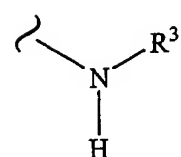
162



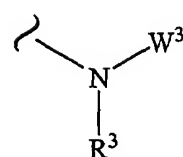
163



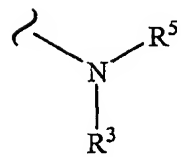
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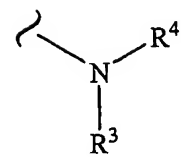
165



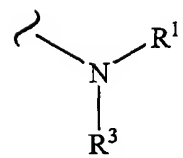
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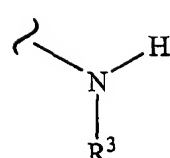
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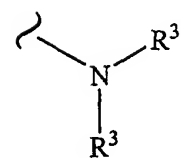
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169

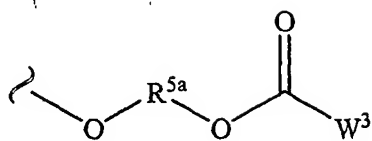


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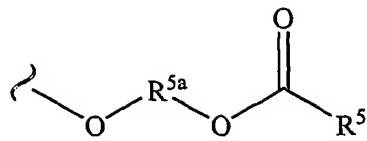


171

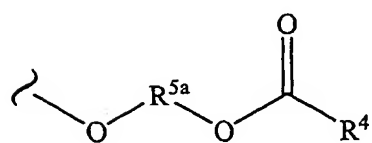
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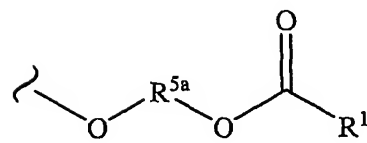
172



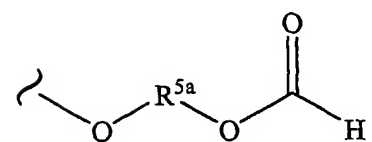
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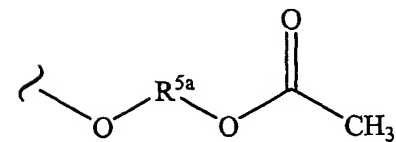
174



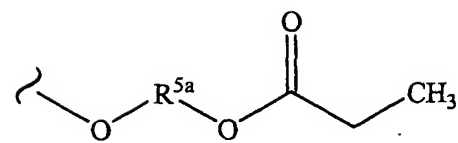
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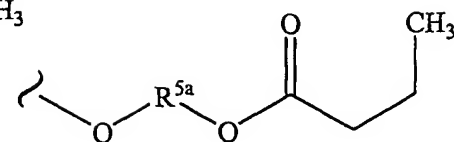
176



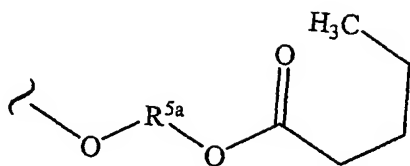
177



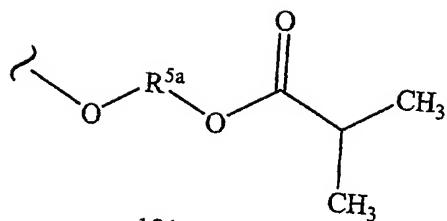
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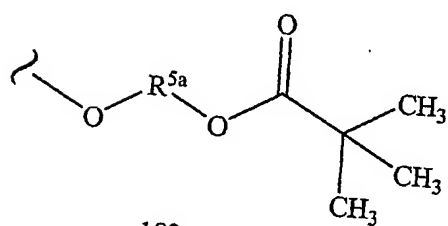
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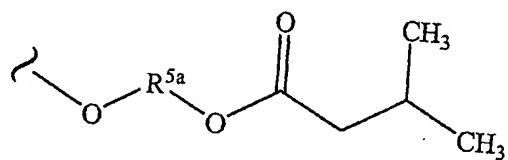
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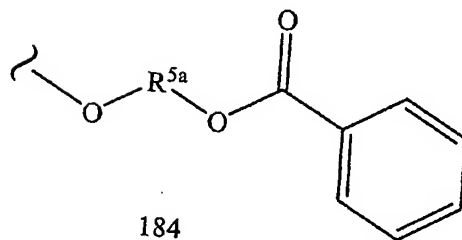
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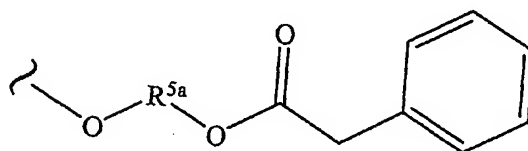
182



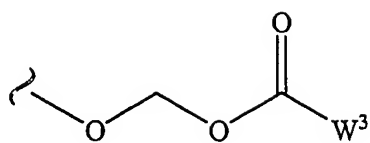
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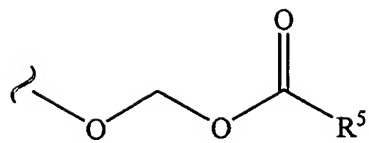
184



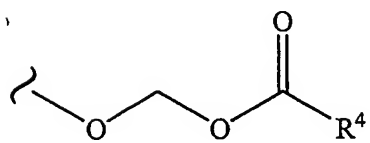
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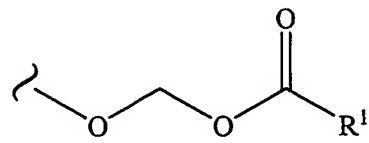
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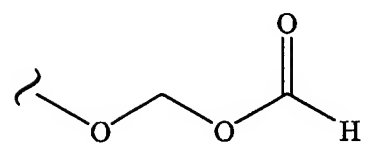
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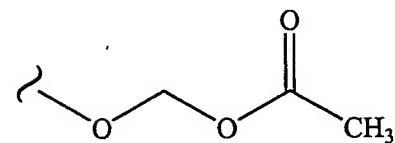
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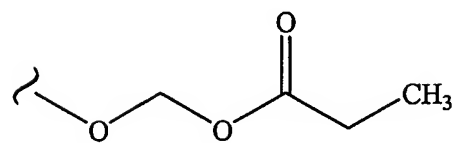
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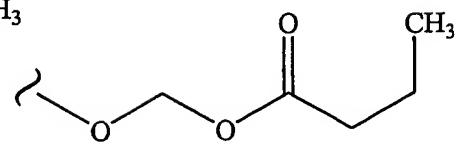
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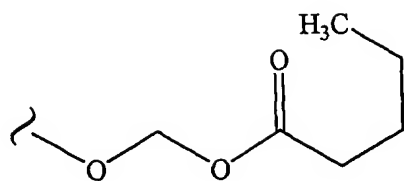


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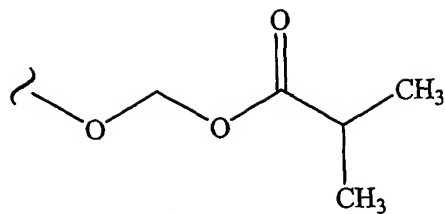


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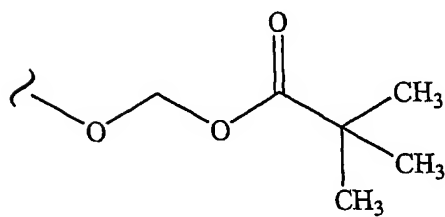
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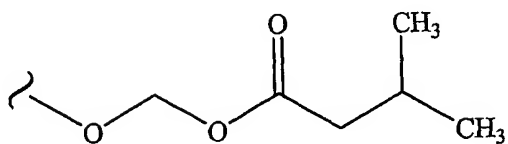
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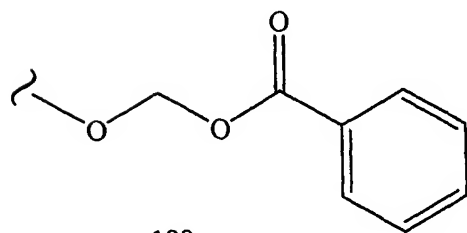
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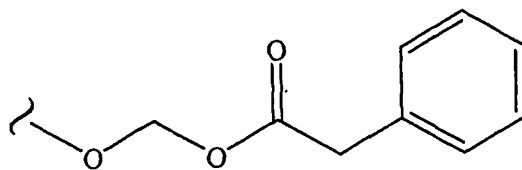
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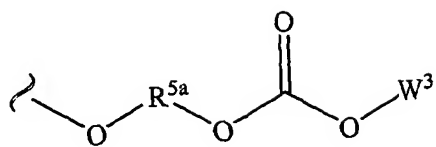
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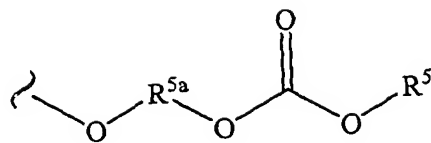
198



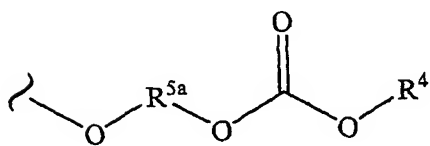
199

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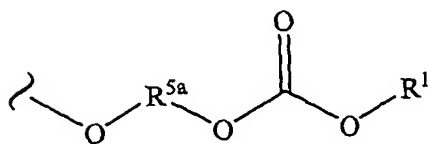
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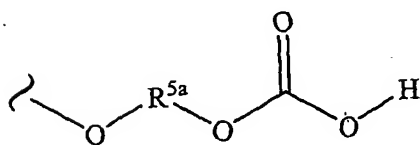
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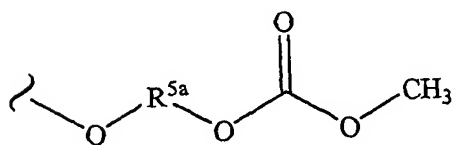
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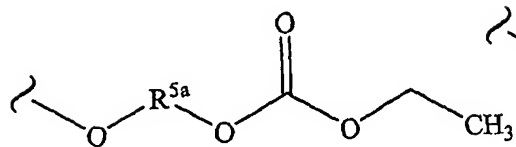
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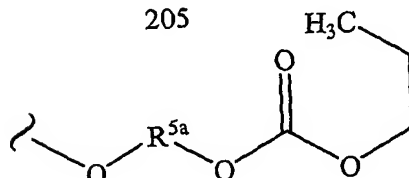
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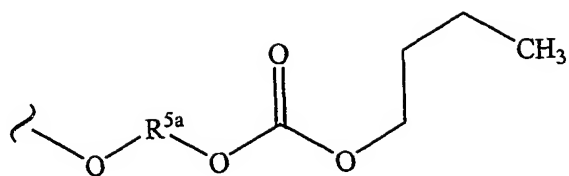
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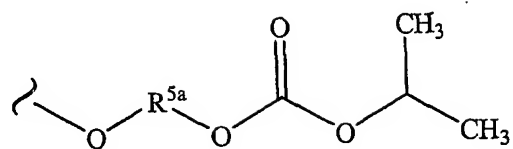
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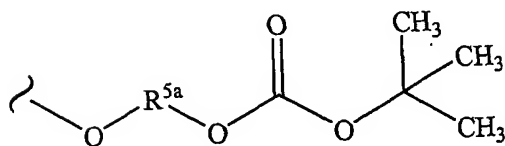
207

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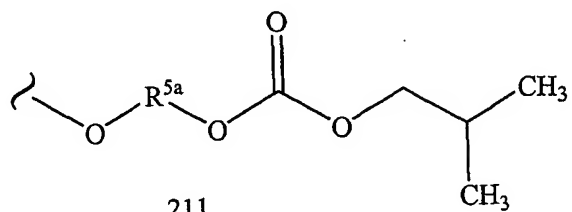
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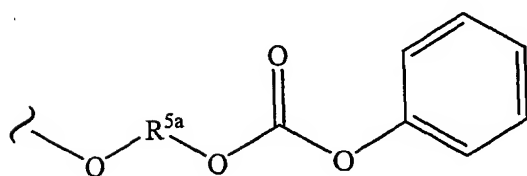
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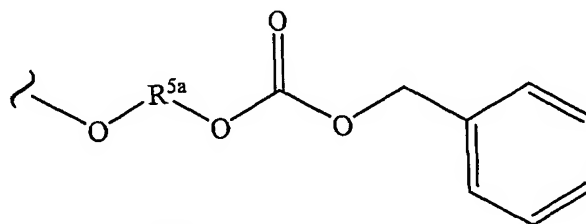
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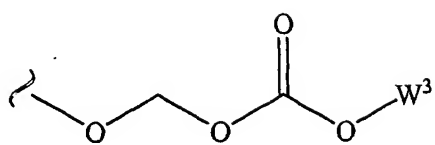
211



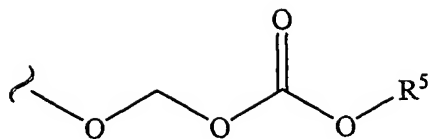
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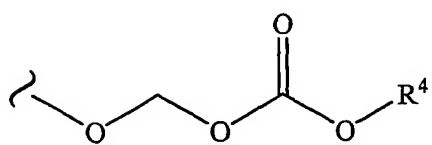
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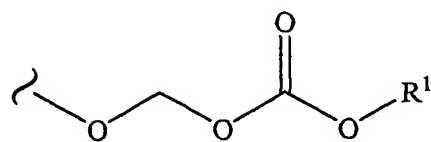
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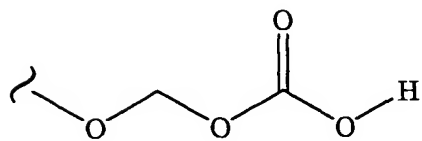
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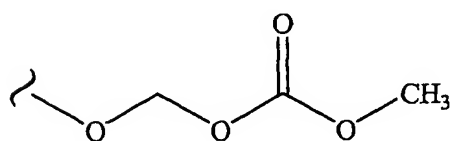
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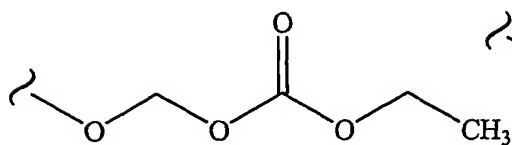
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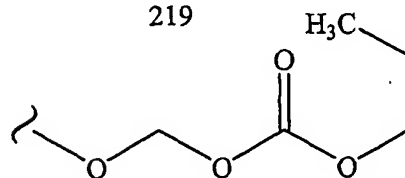
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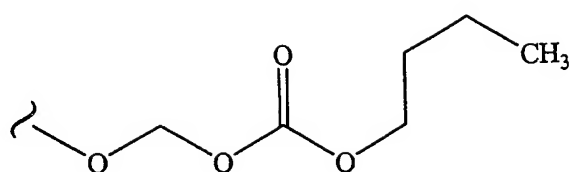


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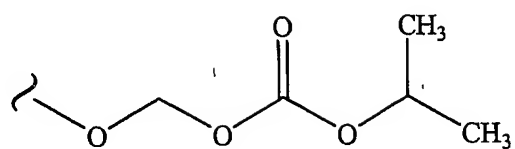


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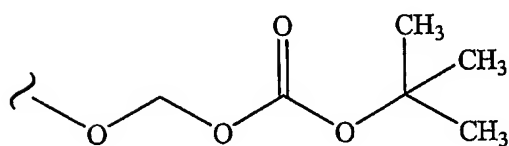
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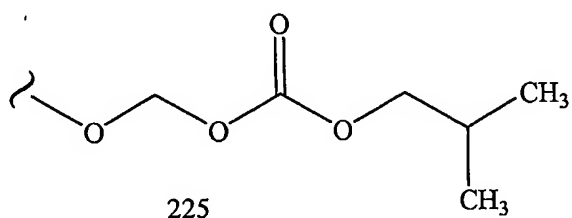
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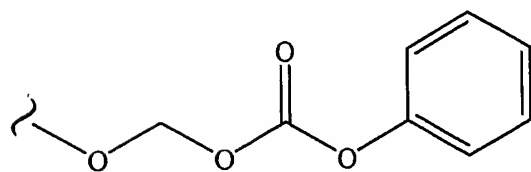
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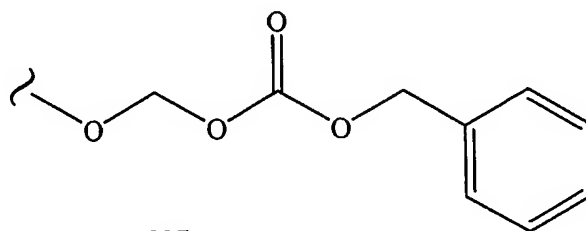
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225



226



227

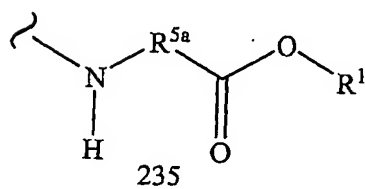
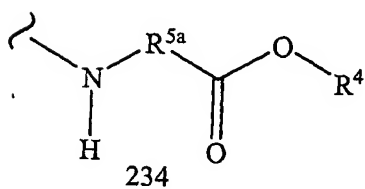
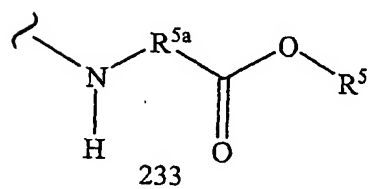
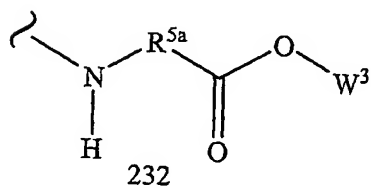
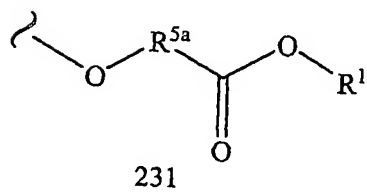
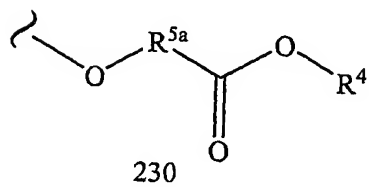
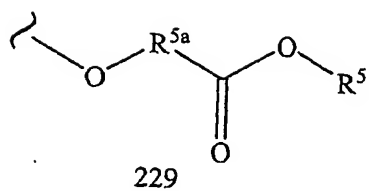
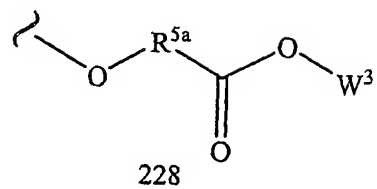
Table 20.35

Table 20.36

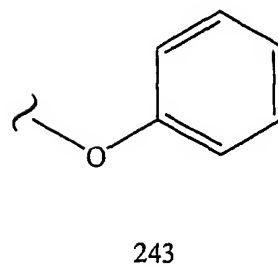
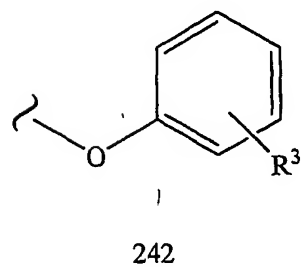
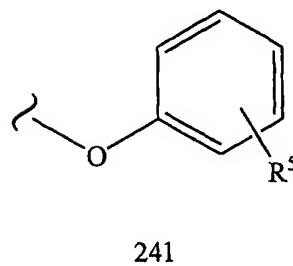
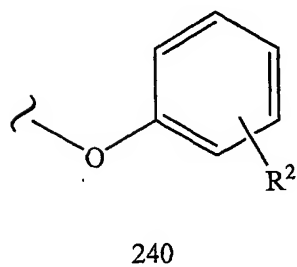
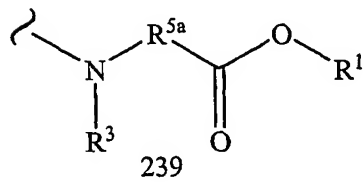
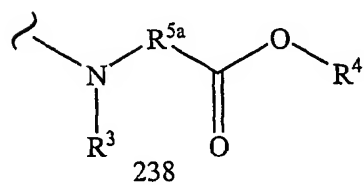
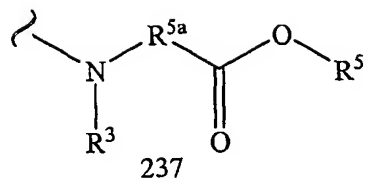
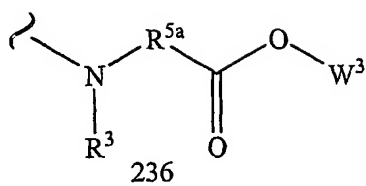


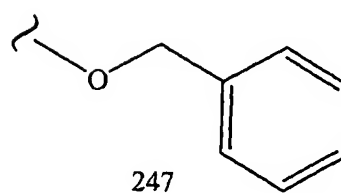
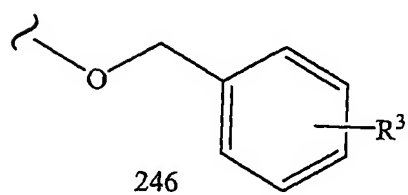
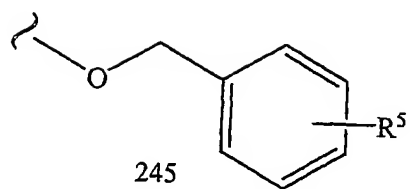
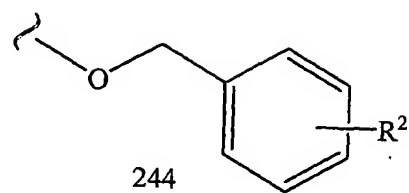
Table 20.37

Table 100

Prodrugs of 1.B

- 1.B.228.228; 1.B.228.229; 1.B.228.230; 1.B.228.231; 1.B.228.236;
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1.B.228.169; 1.B.228.172; 1.B.228.175; 1.B.228.240; 1.B.228.244; 1.B.229.228;
1.B.229.229; 1.B.229.230; 1.B.229.231; 1.B.229.236; 1.B.229.237; 1.B.229.238;
1.B.229.239; 1.B.229.154; 1.B.229.157; 1.B.229.166; 1.B.229.169; 1.B.229.172;
1.B.229.175; 1.B.229.240; 1.B.229.244; 1.B.230.228; 1.B.230.229; 1.B.230.230;
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1.B.230.157; 1.B.230.166; 1.B.230.169; 1.B.230.172; 1.B.230.175; 1.B.230.240;
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1.B.231.169; 1.B.231.172; 1.B.231.175; 1.B.231.240; 1.B.231.244; 1.B.236.228;
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1.B.236.239; 1.B.236.154; 1.B.236.157; 1.B.236.166; 1.B.236.169; 1.B.236.172;
1.B.236.175; 1.B.236.240; 1.B.236.244; 1.B.237.228; 1.B.237.229; 1.B.237.230;
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1.B.237.157; 1.B.237.166; 1.B.237.169; 1.B.237.172; 1.B.237.175; 1.B.237.240;
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1.B.238.237; 1.B.238.238; 1.B.238.239; 1.B.238.154; 1.B.238.157; 1.B.238.166;
1.B.238.169; 1.B.238.172; 1.B.238.175; 1.B.238.240; 1.B.238.244; 1.B.239.228;
1.B.239.229; 1.B.239.230; 1.B.239.231; 1.B.239.236; 1.B.239.237; 1.B.239.238;
1.B.239.239; 1.B.239.154; 1.B.239.157; 1.B.239.166; 1.B.239.169; 1.B.239.172;
25 1.B.239.175; 1.B.239.240; 1.B.239.244; 1.B.154.228; 1.B.154.229; 1.B.154.230;
1.B.154.231; 1.B.154.236; 1.B.154.237; 1.B.154.238; 1.B.154.239; 1.B.154.154;
1.B.154.157; 1.B.154.166; 1.B.154.169; 1.B.154.172; 1.B.154.175; 1.B.154.240;
1.B.154.244; 1.B.157.228; 1.B.157.229; 1.B.157.230; 1.B.157.231; 1.B.157.236;
1.B.157.237; 1.B.157.238; 1.B.157.239; 1.B.157.154; 1.B.157.157; 1.B.157.166;
30 1.B.157.169; 1.B.157.172; 1.B.157.175; 1.B.157.240; 1.B.157.244; 1.B.166.228;

1.B.166.229; 1.B.166.230; 1.B.166.231; 1.B.166.236; 1.B.166.237; 1.B.166.238;
1.B.166.239; 1.B.166.154; 1.B.166.157; 1.B.166.166; 1.B.166.169; 1.B.166.172;
1.B.166.175; 1.B.166.240; 1.B.166.244; 1.B.169.228; 1.B.169.229; 1.B.169.230;
1.B.169.231; 1.B.169.236; 1.B.169.237; 1.B.169.238; 1.B.169.239; 1.B.169.154;
5 1.B.169.157; 1.B.169.166; 1.B.169.169; 1.B.169.172; 1.B.169.175; 1.B.169.240;
1.B.169.244; 1.B.172.228; 1.B.172.229; 1.B.172.230; 1.B.172.231; 1.B.172.236;
1.B.172.237; 1.B.172.238; 1.B.172.239; 1.B.172.154; 1.B.172.157; 1.B.172.166;
1.B.172.169; 1.B.172.172; 1.B.172.175; 1.B.172.240; 1.B.172.244; 1.B.175.228;
1.B.175.229; 1.B.175.230; 1.B.175.231; 1.B.175.236; 1.B.175.237; 1.B.175.238;
10 1.B.175.239; 1.B.175.154; 1.B.175.157; 1.B.175.166; 1.B.175.169; 1.B.175.172;
1.B.175.175; 1.B.175.240; 1.B.175.244; 1.B.240.228; 1.B.240.229; 1.B.240.230;
1.B.240.231; 1.B.240.236; 1.B.240.237; 1.B.240.238; 1.B.240.239; 1.B.240.154;
1.B.240.157; 1.B.240.166; 1.B.240.169; 1.B.240.172; 1.B.240.175; 1.B.240.240;
1.B.240.244; 1.B.244.228; 1.B.244.229; 1.B.244.230; 1.B.244.231; 1.B.244.236;
15 1.B.244.237; 1.B.244.238; 1.B.244.239; 1.B.244.154; 1.B.244.157; 1.B.244.166;
1.B.244.169; 1.B.244.172; 1.B.244.175; 1.B.244.240; 1.B.244.244;

Prodrugs of 1.D

1.D.228.228; 1.D.228.229; 1.D.228.230; 1.D.228.231; 1.D.228.236;
20 1.D.228.237; 1.D.228.238; 1.D.228.239; 1.D.228.154; 1.D.228.157;
1.D.228.166; 1.D.228.169; 1.D.228.172; 1.D.228.175; 1.D.228.240;
1.D.228.244; 1.D.229.228; 1.D.229.229; 1.D.229.230; 1.D.229.231;
1.D.229.236; 1.D.229.237; 1.D.229.238; 1.D.229.239; 1.D.229.154;
1.D.229.157; 1.D.229.166; 1.D.229.169; 1.D.229.172; 1.D.229.175;
25 1.D.229.240; 1.D.229.244; 1.D.230.228; 1.D.230.229; 1.D.230.230;
1.D.230.231; 1.D.230.236; 1.D.230.237; 1.D.230.238; 1.D.230.239;
1.D.230.154; 1.D.230.157; 1.D.230.166; 1.D.230.169; 1.D.230.172;
1.D.230.175; 1.D.230.240; 1.D.230.244; 1.D.231.228; 1.D.231.229;
1.D.231.230; 1.D.231.231; 1.D.231.236; 1.D.231.237; 1.D.231.238;
30 1.D.231.239; 1.D.231.154; 1.D.231.157; 1.D.231.166; 1.D.231.169;

- 1.D.231.172; 1.D.231.175; 1.D.231.240; 1.D.231.244; 1.D.236.228;
1.D.236.229; 1.D.236.230; 1.D.236.231; 1.D.236.236; 1.D.236.237;
1.D.236.238; 1.D.236.239; 1.D.236.154; 1.D.236.157; 1.D.236.166;
1.D.236.169; 1.D.236.172; 1.D.236.175; 1.D.236.240; 1.D.236.244;
5 1.D.237.228; 1.D.237.229; 1.D.237.230; 1.D.237.231; 1.D.237.236;
1.D.237.237; 1.D.237.238; 1.D.237.239; 1.D.237.154; 1.D.237.157;
1.D.237.166; 1.D.237.169; 1.D.237.172; 1.D.237.175; 1.D.237.240;
1.D.237.244; 1.D.238.228; 1.D.238.229; 1.D.238.230; 1.D.238.231;
1.D.238.236; 1.D.238.237; 1.D.238.238; 1.D.238.239; 1.D.238.154;
10 1.D.238.157; 1.D.238.166; 1.D.238.169; 1.D.238.172; 1.D.238.175;
1.D.238.240; 1.D.238.244; 1.D.239.228; 1.D.239.229; 1.D.239.230;
1.D.239.231; 1.D.239.236; 1.D.239.237; 1.D.239.238; 1.D.239.239;
1.D.239.154; 1.D.239.157; 1.D.239.166; 1.D.239.169; 1.D.239.172;
1.D.239.175; 1.D.239.240; 1.D.239.244; 1.D.154.228; 1.D.154.229;
15 1.D.154.230; 1.D.154.231; 1.D.154.236; 1.D.154.237; 1.D.154.238;
1.D.154.239; 1.D.154.154; 1.D.154.157; 1.D.154.166; 1.D.154.169;
1.D.154.172; 1.D.154.175; 1.D.154.240; 1.D.154.244; 1.D.157.228;
1.D.157.229; 1.D.157.230; 1.D.157.231; 1.D.157.236; 1.D.157.237;
1.D.157.238; 1.D.157.239; 1.D.157.154; 1.D.157.157; 1.D.157.166;
20 1.D.157.169; 1.D.157.172; 1.D.157.175; 1.D.157.240; 1.D.157.244;
1.D.166.228; 1.D.166.229; 1.D.166.230; 1.D.166.231; 1.D.166.236;
1.D.166.237; 1.D.166.238; 1.D.166.239; 1.D.166.154; 1.D.166.157;
1.D.166.166; 1.D.166.169; 1.D.166.172; 1.D.166.175; 1.D.166.240;
1.D.166.244; 1.D.169.228; 1.D.169.229; 1.D.169.230; 1.D.169.231;
25 1.D.169.236; 1.D.169.237; 1.D.169.238; 1.D.169.239; 1.D.169.154;
1.D.169.157; 1.D.169.166; 1.D.169.169; 1.D.169.172; 1.D.169.175;
1.D.169.240; 1.D.169.244; 1.D.172.228; 1.D.172.229; 1.D.172.230;
1.D.172.231; 1.D.172.236; 1.D.172.237; 1.D.172.238; 1.D.172.239;
1.D.172.154; 1.D.172.157; 1.D.172.166; 1.D.172.169; 1.D.172.172;
30 1.D.172.175; 1.D.172.240; 1.D.172.244; 1.D.175.228; 1.D.175.229;

1.D.175.230; 1.D.175.231; 1.D.175.236; 1.D.175.237; 1.D.175.238;
 1.D.175.239; 1.D.175.154; 1.D.175.157; 1.D.175.166; 1.D.175.169;
 1.D.175.172; 1.D.175.175; 1.D.175.240; 1.D.175.244; 1.D.240.228;
 1.D.240.229; 1.D.240.230; 1.D.240.231; 1.D.240.236; 1.D.240.237;
 5 1.D.240.238; 1.D.240.239; 1.D.240.154; 1.D.240.157; 1.D.240.166;
 1.D.240.169; 1.D.240.172; 1.D.240.175; 1.D.240.240; 1.D.240.244;
 1.D.244.228; 1.D.244.229; 1.D.244.230; 1.D.244.231; 1.D.244.236;
 1.D.244.237; 1.D.244.238; 1.D.244.239; 1.D.244.154; 1.D.244.157;
 1.D.244.166; 1.D.244.169; 1.D.244.172; 1.D.244.175; 1.D.244.240;
 10 1.D.244.244;

Prodrugs of 1.E

1.E.228.228; 1.E.228.229; 1.E.228.230; 1.E.228.231; 1.E.228.236;
 1.E.228.237; 1.E.228.238; 1.E.228.239; 1.E.228.154; 1.E.228.157; 1.E.228.166;
 15 1.E.228.169; 1.E.228.172; 1.E.228.175; 1.E.228.240; 1.E.228.244; 1.E.229.228;
 1.E.229.229; 1.E.229.230; 1.E.229.231; 1.E.229.236; 1.E.229.237; 1.E.229.238;
 1.E.229.239; 1.E.229.154; 1.E.229.157; 1.E.229.166; 1.E.229.169; 1.E.229.172;
 1.E.229.175; 1.E.229.240; 1.E.229.244; 1.E.230.228; 1.E.230.229; 1.E.230.230;
 1.E.230.231; 1.E.230.236; 1.E.230.237; 1.E.230.238; 1.E.230.239; 1.E.230.154;
 20 1.E.230.157; 1.E.230.166; 1.E.230.169; 1.E.230.172; 1.E.230.175; 1.E.230.240;
 1.E.230.244; 1.E.231.228; 1.E.231.229; 1.E.231.230; 1.E.231.231; 1.E.231.236;
 1.E.231.237; 1.E.231.238; 1.E.231.239; 1.E.231.154; 1.E.231.157; 1.E.231.166;
 1.E.231.169; 1.E.231.172; 1.E.231.175; 1.E.231.240; 1.E.231.244; 1.E.236.228;
 1.E.236.229; 1.E.236.230; 1.E.236.231; 1.E.236.236; 1.E.236.237; 1.E.236.238;
 25 1.E.236.239; 1.E.236.154; 1.E.236.157; 1.E.236.166; 1.E.236.169; 1.E.236.172;
 1.E.236.175; 1.E.236.240; 1.E.236.244; 1.E.237.228; 1.E.237.229; 1.E.237.230;
 1.E.237.231; 1.E.237.236; 1.E.237.237; 1.E.237.238; 1.E.237.239; 1.E.237.154;
 1.E.237.157; 1.E.237.166; 1.E.237.169; 1.E.237.172; 1.E.237.175; 1.E.237.240;
 1.E.237.244; 1.E.238.228; 1.E.238.229; 1.E.238.230; 1.E.238.231; 1.E.238.236;
 30 1.E.238.237; 1.E.238.238; 1.E.238.239; 1.E.238.154; 1.E.238.157; 1.E.238.166;

1.E.238.169; 1.E.238.172; 1.E.238.175; 1.E.238.240; 1.E.238.244; 1.E.239.228;
1.E.239.229; 1.E.239.230; 1.E.239.231; 1.E.239.236; 1.E.239.237; 1.E.239.238;
1.E.239.239; 1.E.239.154; 1.E.239.157; 1.E.239.166; 1.E.239.169; 1.E.239.172;
1.E.239.175; 1.E.239.240; 1.E.239.244; 1.E.154.228; 1.E.154.229; 1.E.154.230;
5 1.E.154.231; 1.E.154.236; 1.E.154.237; 1.E.154.238; 1.E.154.239; 1.E.154.154;
1.E.154.157; 1.E.154.166; 1.E.154.169; 1.E.154.172; 1.E.154.175; 1.E.154.240;
1.E.154.244; 1.E.157.228; 1.E.157.229; 1.E.157.230; 1.E.157.231; 1.E.157.236;
1.E.157.237; 1.E.157.238; 1.E.157.239; 1.E.157.154; 1.E.157.157; 1.E.157.166;
1.E.157.169; 1.E.157.172; 1.E.157.175; 1.E.157.240; 1.E.157.244; 1.E.166.228;
10 1.E.166.229; 1.E.166.230; 1.E.166.231; 1.E.166.236; 1.E.166.237; 1.E.166.238;
1.E.166.239; 1.E.166.154; 1.E.166.157; 1.E.166.166; 1.E.166.169; 1.E.166.172;
1.E.166.175; 1.E.166.240; 1.E.166.244; 1.E.169.228; 1.E.169.229; 1.E.169.230;
1.E.169.231; 1.E.169.236; 1.E.169.237; 1.E.169.238; 1.E.169.239; 1.E.169.154;
1.E.169.157; 1.E.169.166; 1.E.169.169; 1.E.169.172; 1.E.169.175; 1.E.169.240;
15 1.E.169.244; 1.E.172.228; 1.E.172.229; 1.E.172.230; 1.E.172.231; 1.E.172.236;
1.E.172.237; 1.E.172.238; 1.E.172.239; 1.E.172.154; 1.E.172.157; 1.E.172.166;
1.E.172.169; 1.E.172.172; 1.E.172.175; 1.E.172.240; 1.E.172.244; 1.E.175.228;
1.E.175.229; 1.E.175.230; 1.E.175.231; 1.E.175.236; 1.E.175.237; 1.E.175.238;
1.E.175.239; 1.E.175.154; 1.E.175.157; 1.E.175.166; 1.E.175.169; 1.E.175.172;
20 1.E.175.175; 1.E.175.240; 1.E.175.244; 1.E.240.228; 1.E.240.229; 1.E.240.230;
1.E.240.231; 1.E.240.236; 1.E.240.237; 1.E.240.238; 1.E.240.239; 1.E.240.154;
1.E.240.157; 1.E.240.166; 1.E.240.169; 1.E.240.172; 1.E.240.175; 1.E.240.240;
1.E.240.244; 1.E.244.228; 1.E.244.229; 1.E.244.230; 1.E.244.231; 1.E.244.236;
1.E.244.237; 1.E.244.238; 1.E.244.239; 1.E.244.154; 1.E.244.157; 1.E.244.166;
25 1.E.244.169; 1.E.244.172; 1.E.244.175; 1.E.244.240; 1.E.244.244;

Prodrugs of 1.G

1.G.228.228; 1.G.228.229; 1.G.228.230; 1.G.228.231; 1.G.228.236;
1.G.228.237; 1.G.228.238; 1.G.228.239; 1.G.228.154; 1.G.228.157;
30 1.G.228.166; 1.G.228.169; 1.G.228.172; 1.G.228.175; 1.G.228.240;

- 1.G.228.244; 1.G.229.228; 1.G.229.229; 1.G.229.230; 1.G.229.231;
1.G.229.236; 1.G.229.237; 1.G.229.238; 1.G.229.239; 1.G.229.154;
1.G.229.157; 1.G.229.166; 1.G.229.169; 1.G.229.172; 1.G.229.175;
1.G.229.240; 1.G.229.244; 1.G.230.228; 1.G.230.229; 1.G.230.230;
5 1.G.230.231; 1.G.230.236; 1.G.230.237; 1.G.230.238; 1.G.230.239;
1.G.230.154; 1.G.230.157; 1.G.230.166; 1.G.230.169; 1.G.230.172;
1.G.230.175; 1.G.230.240; 1.G.230.244; 1.G.231.228; 1.G.231.229;
1.G.231.230; 1.G.231.231; 1.G.231.236; 1.G.231.237; 1.G.231.238;
1.G.231.239; 1.G.231.154; 1.G.231.157; 1.G.231.166; 1.G.231.169;
10 1.G.231.172; 1.G.231.175; 1.G.231.240; 1.G.231.244; 1.G.236.228;
1.G.236.229; 1.G.236.230; 1.G.236.231; 1.G.236.236; 1.G.236.237;
1.G.236.238; 1.G.236.239; 1.G.236.154; 1.G.236.157; 1.G.236.166;
1.G.236.169; 1.G.236.172; 1.G.236.175; 1.G.236.240; 1.G.236.244;
1.G.237.228; 1.G.237.229; 1.G.237.230; 1.G.237.231; 1.G.237.236;
15 1.G.237.237; 1.G.237.238; 1.G.237.239; 1.G.237.154; 1.G.237.157;
1.G.237.166; 1.G.237.169; 1.G.237.172; 1.G.237.175; 1.G.237.240;
1.G.237.244; 1.G.238.228; 1.G.238.229; 1.G.238.230; 1.G.238.231;
1.G.238.236; 1.G.238.237; 1.G.238.238; 1.G.238.239; 1.G.238.154;
1.G.238.157; 1.G.238.166; 1.G.238.169; 1.G.238.172; 1.G.238.175;
20 1.G.238.240; 1.G.238.244; 1.G.239.228; 1.G.239.229; 1.G.239.230;
1.G.239.231; 1.G.239.236; 1.G.239.237; 1.G.239.238; 1.G.239.239;
1.G.239.154; 1.G.239.157; 1.G.239.166; 1.G.239.169; 1.G.239.172;
1.G.239.175; 1.G.239.240; 1.G.239.244; 1.G.154.228; 1.G.154.229;
1.G.154.230; 1.G.154.231; 1.G.154.236; 1.G.154.237; 1.G.154.238;
25 1.G.154.239; 1.G.154.154; 1.G.154.157; 1.G.154.166; 1.G.154.169;
1.G.154.172; 1.G.154.175; 1.G.154.240; 1.G.154.244; 1.G.157.228;
1.G.157.229; 1.G.157.230; 1.G.157.231; 1.G.157.236; 1.G.157.237;
1.G.157.238; 1.G.157.239; 1.G.157.154; 1.G.157.157; 1.G.157.166;
1.G.157.169; 1.G.157.172; 1.G.157.175; 1.G.157.240; 1.G.157.244;
30 1.G.166.228; 1.G.166.229; 1.G.166.230; 1.G.166.231; 1.G.166.236;

1.G.166.237; 1.G.166.238; 1.G.166.239; 1.G.166.154; 1.G.166.157;
 1.G.166.166; 1.G.166.169; 1.G.166.172; 1.G.166.175; 1.G.166.240;
 1.G.166.244; 1.G.169.228; 1.G.169.229; 1.G.169.230; 1.G.169.231;
 1.G.169.236; 1.G.169.237; 1.G.169.238; 1.G.169.239; 1.G.169.154;
 5 1.G.169.157; 1.G.169.166; 1.G.169.169; 1.G.169.172; 1.G.169.175;
 1.G.169.240; 1.G.169.244; 1.G.172.228; 1.G.172.229; 1.G.172.230;
 1.G.172.231; 1.G.172.236; 1.G.172.237; 1.G.172.238; 1.G.172.239;
 1.G.172.154; 1.G.172.157; 1.G.172.166; 1.G.172.169; 1.G.172.172;
 1.G.172.175; 1.G.172.240; 1.G.172.244; 1.G.175.228; 1.G.175.229;
 10 1.G.175.230; 1.G.175.231; 1.G.175.236; 1.G.175.237; 1.G.175.238;
 1.G.175.239; 1.G.175.154; 1.G.175.157; 1.G.175.166; 1.G.175.169;
 1.G.175.172; 1.G.175.175; 1.G.175.240; 1.G.175.244; 1.G.240.228;
 1.G.240.229; 1.G.240.230; 1.G.240.231; 1.G.240.236; 1.G.240.237;
 1.G.240.238; 1.G.240.239; 1.G.240.154; 1.G.240.157; 1.G.240.166;
 15 1.G.240.169; 1.G.240.172; 1.G.240.175; 1.G.240.240; 1.G.240.244;
 1.G.244.228; 1.G.244.229; 1.G.244.230; 1.G.244.231; 1.G.244.236;
 1.G.244.237; 1.G.244.238; 1.G.244.239; 1.G.244.154; 1.G.244.157;
 1.G.244.166; 1.G.244.169; 1.G.244.172; 1.G.244.175; 1.G.244.240;
 1.G.244.244;

20

Prodrugs of 1.I

1.I.228.228; 1.I.228.229; 1.I.228.230; 1.I.228.231; 1.I.228.236; 1.I.228.237;
 1.I.228.238; 1.I.228.239; 1.I.228.154; 1.I.228.157; 1.I.228.166; 1.I.228.169;
 1.I.228.172; 1.I.228.175; 1.I.228.240; 1.I.228.244; 1.I.229.228; 1.I.229.229;
 25 1.I.229.230; 1.I.229.231; 1.I.229.236; 1.I.229.237; 1.I.229.238; 1.I.229.239;
 1.I.229.154; 1.I.229.157; 1.I.229.166; 1.I.229.169; 1.I.229.172; 1.I.229.175;
 1.I.229.240; 1.I.229.244; 1.I.230.228; 1.I.230.229; 1.I.230.230; 1.I.230.231;
 1.I.230.236; 1.I.230.237; 1.I.230.238; 1.I.230.239; 1.I.230.154; 1.I.230.157;
 1.I.230.166; 1.I.230.169; 1.I.230.172; 1.I.230.175; 1.I.230.240; 1.I.230.244;
 30 1.I.231.228; 1.I.231.229; 1.I.231.230; 1.I.231.231; 1.I.231.236; 1.I.231.237;

1.I.231.238; 1.I.231.239; 1.I.231.154; 1.I.231.157; 1.I.231.166; 1.I.231.169;
1.I.231.172; 1.I.231.175; 1.I.231.240; 1.I.231.244; 1.I.236.228; 1.I.236.229;
1.I.236.230; 1.I.236.231; 1.I.236.236; 1.I.236.237; 1.I.236.238; 1.I.236.239;
1.I.236.154; 1.I.236.157; 1.I.236.166; 1.I.236.169; 1.I.236.172; 1.I.236.175;
5 1.I.236.240; 1.I.236.244; 1.I.237.228; 1.I.237.229; 1.I.237.230; 1.I.237.231;
1.I.237.236; 1.I.237.237; 1.I.237.238; 1.I.237.239; 1.I.237.154; 1.I.237.157;
1.I.237.166; 1.I.237.169; 1.I.237.172; 1.I.237.175; 1.I.237.240; 1.I.237.244;
1.I.238.228; 1.I.238.229; 1.I.238.230; 1.I.238.231; 1.I.238.236; 1.I.238.237;
1.I.238.238; 1.I.238.239; 1.I.238.154; 1.I.238.157; 1.I.238.166; 1.I.238.169;
10 1.I.238.172; 1.I.238.175; 1.I.238.240; 1.I.238.244; 1.I.239.228; 1.I.239.229;
1.I.239.230; 1.I.239.231; 1.I.239.236; 1.I.239.237; 1.I.239.238; 1.I.239.239;
1.I.239.154; 1.I.239.157; 1.I.239.166; 1.I.239.169; 1.I.239.172; 1.I.239.175;
1.I.239.240; 1.I.239.244; 1.I.154.228; 1.I.154.229; 1.I.154.230; 1.I.154.231;
1.I.154.236; 1.I.154.237; 1.I.154.238; 1.I.154.239; 1.I.154.154; 1.I.154.157;
15 1.I.154.166; 1.I.154.169; 1.I.154.172; 1.I.154.175; 1.I.154.240; 1.I.154.244;
1.I.157.228; 1.I.157.229; 1.I.157.230; 1.I.157.231; 1.I.157.236; 1.I.157.237;
1.I.157.238; 1.I.157.239; 1.I.157.154; 1.I.157.157; 1.I.157.166; 1.I.157.169;
1.I.157.172; 1.I.157.175; 1.I.157.240; 1.I.157.244; 1.I.166.228; 1.I.166.229;
1.I.166.230; 1.I.166.231; 1.I.166.236; 1.I.166.237; 1.I.166.238; 1.I.166.239;
20 1.I.166.154; 1.I.166.157; 1.I.166.166; 1.I.166.169; 1.I.166.172; 1.I.166.175;
1.I.166.240; 1.I.166.244; 1.I.169.228; 1.I.169.229; 1.I.169.230; 1.I.169.231;
1.I.169.236; 1.I.169.237; 1.I.169.238; 1.I.169.239; 1.I.169.154; 1.I.169.157;
1.I.169.166; 1.I.169.169; 1.I.169.172; 1.I.169.175; 1.I.169.240; 1.I.169.244;
1.I.172.228; 1.I.172.229; 1.I.172.230; 1.I.172.231; 1.I.172.236; 1.I.172.237;
25 1.I.172.238; 1.I.172.239; 1.I.172.154; 1.I.172.157; 1.I.172.166; 1.I.172.169;
1.I.172.172; 1.I.172.175; 1.I.172.240; 1.I.172.244; 1.I.175.228; 1.I.175.229;
1.I.175.230; 1.I.175.231; 1.I.175.236; 1.I.175.237; 1.I.175.238; 1.I.175.239;
1.I.175.154; 1.I.175.157; 1.I.175.166; 1.I.175.169; 1.I.175.172; 1.I.175.175;
1.I.175.240; 1.I.175.244; 1.I.240.228; 1.I.240.229; 1.I.240.230; 1.I.240.231;
30 1.I.240.236; 1.I.240.237; 1.I.240.238; 1.I.240.239; 1.I.240.154; 1.I.240.157;

1.I.240.166; 1.I.240.169; 1.I.240.172; 1.I.240.175; 1.I.240.240; 1.I.240.244;
1.I.244.228; 1.I.244.229; 1.I.244.230; 1.I.244.231; 1.I.244.236; 1.I.244.237;
1.I.244.238; 1.I.244.239; 1.I.244.154; 1.I.244.157; 1.I.244.166; 1.I.244.169;
1.I.244.172; 1.I.244.175; 1.I.244.240; 1.I.244.244;

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Prodrugs of 1.J

1.J.228.228; 1.J.228.229; 1.J.228.230; 1.J.228.231; 1.J.228.236; 1.J.228.237;
1.J.228.238; 1.J.228.239; 1.J.228.154; 1.J.228.157; 1.J.228.166; 1.J.228.169;
1.J.228.172; 1.J.228.175; 1.J.228.240; 1.J.228.244; 1.J.229.228; 1.J.229.229;
10 1.J.229.230; 1.J.229.231; 1.J.229.236; 1.J.229.237; 1.J.229.238; 1.J.229.239;
1.J.229.154; 1.J.229.157; 1.J.229.166; 1.J.229.169; 1.J.229.172; 1.J.229.175;
1.J.229.240; 1.J.229.244; 1.J.230.228; 1.J.230.229; 1.J.230.230; 1.J.230.231;
1.J.230.236; 1.J.230.237; 1.J.230.238; 1.J.230.239; 1.J.230.154; 1.J.230.157;
1.J.230.166; 1.J.230.169; 1.J.230.172; 1.J.230.175; 1.J.230.240; 1.J.230.244;
15 1.J.231.228; 1.J.231.229; 1.J.231.230; 1.J.231.231; 1.J.231.236; 1.J.231.237;
1.J.231.238; 1.J.231.239; 1.J.231.154; 1.J.231.157; 1.J.231.166; 1.J.231.169;
1.J.231.172; 1.J.231.175; 1.J.231.240; 1.J.231.244; 1.J.236.228; 1.J.236.229;
1.J.236.230; 1.J.236.231; 1.J.236.236; 1.J.236.237; 1.J.236.238; 1.J.236.239;
1.J.236.154; 1.J.236.157; 1.J.236.166; 1.J.236.169; 1.J.236.172; 1.J.236.175;
20 1.J.236.240; 1.J.236.244; 1.J.237.228; 1.J.237.229; 1.J.237.230; 1.J.237.231;
1.J.237.236; 1.J.237.237; 1.J.237.238; 1.J.237.239; 1.J.237.154; 1.J.237.157;
1.J.237.166; 1.J.237.169; 1.J.237.172; 1.J.237.175; 1.J.237.240; 1.J.237.244;
1.J.238.228; 1.J.238.229; 1.J.238.230; 1.J.238.231; 1.J.238.236; 1.J.238.237;
1.J.238.238; 1.J.238.239; 1.J.238.154; 1.J.238.157; 1.J.238.166; 1.J.238.169;
25 1.J.238.172; 1.J.238.175; 1.J.238.240; 1.J.238.244; 1.J.239.228; 1.J.239.229;
1.J.239.230; 1.J.239.231; 1.J.239.236; 1.J.239.237; 1.J.239.238; 1.J.239.239;
1.J.239.154; 1.J.239.157; 1.J.239.166; 1.J.239.169; 1.J.239.172; 1.J.239.175;
1.J.239.240; 1.J.239.244; 1.J.154.228; 1.J.154.229; 1.J.154.230; 1.J.154.231;
1.J.154.236; 1.J.154.237; 1.J.154.238; 1.J.154.239; 1.J.154.154; 1.J.154.157;
30 1.J.154.166; 1.J.154.169; 1.J.154.172; 1.J.154.175; 1.J.154.240; 1.J.154.244;

1.J.157.228; 1.J.157.229; 1.J.157.230; 1.J.157.231; 1.J.157.236; 1.J.157.237;
 1.J.157.238; 1.J.157.239; 1.J.157.154; 1.J.157.157; 1.J.157.166; 1.J.157.169;
 1.J.157.172; 1.J.157.175; 1.J.157.240; 1.J.157.244; 1.J.166.228; 1.J.166.229;
 1.J.166.230; 1.J.166.231; 1.J.166.236; 1.J.166.237; 1.J.166.238; 1.J.166.239;
 5 1.J.166.154; 1.J.166.157; 1.J.166.166; 1.J.166.169; 1.J.166.172; 1.J.166.175;
 1.J.166.240; 1.J.166.244; 1.J.169.228; 1.J.169.229; 1.J.169.230; 1.J.169.231;
 1.J.169.236; 1.J.169.237; 1.J.169.238; 1.J.169.239; 1.J.169.154; 1.J.169.157;
 1.J.169.166; 1.J.169.169; 1.J.169.172; 1.J.169.175; 1.J.169.240; 1.J.169.244;
 1.J.172.228; 1.J.172.229; 1.J.172.230; 1.J.172.231; 1.J.172.236; 1.J.172.237;
 10 1.J.172.238; 1.J.172.239; 1.J.172.154; 1.J.172.157; 1.J.172.166; 1.J.172.169;
 1.J.172.172; 1.J.172.175; 1.J.172.240; 1.J.172.244; 1.J.175.228; 1.J.175.229;
 1.J.175.230; 1.J.175.231; 1.J.175.236; 1.J.175.237; 1.J.175.238; 1.J.175.239;
 1.J.175.154; 1.J.175.157; 1.J.175.166; 1.J.175.169; 1.J.175.172; 1.J.175.175;
 1.J.175.240; 1.J.175.244; 1.J.240.228; 1.J.240.229; 1.J.240.230; 1.J.240.231;
 15 1.J.240.236; 1.J.240.237; 1.J.240.238; 1.J.240.239; 1.J.240.154; 1.J.240.157;
 1.J.240.166; 1.J.240.169; 1.J.240.172; 1.J.240.175; 1.J.240.240; 1.J.240.244;
 1.J.244.228; 1.J.244.229; 1.J.244.230; 1.J.244.231; 1.J.244.236; 1.J.244.237;
 1.J.244.238; 1.J.244.239; 1.J.244.154; 1.J.244.157; 1.J.244.166; 1.J.244.169;
 1.J.244.172; 1.J.244.175; 1.J.244.240; 1.J.244.244;

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Prodrugs of 1.L

1.L.228.228; 1.L.228.229; 1.L.228.230; 1.L.228.231; 1.L.228.236;
 1.L.228.237; 1.L.228.238; 1.L.228.239; 1.L.228.154; 1.L.228.157; 1.L.228.166;
 1.L.228.169; 1.L.228.172; 1.L.228.175; 1.L.228.240; 1.L.228.244; 1.L.229.228;
 25 1.L.229.229; 1.L.229.230; 1.L.229.231; 1.L.229.236; 1.L.229.237; 1.L.229.238;
 1.L.229.239; 1.L.229.154; 1.L.229.157; 1.L.229.166; 1.L.229.169; 1.L.229.172;
 1.L.229.175; 1.L.229.240; 1.L.229.244; 1.L.230.228; 1.L.230.229; 1.L.230.230;
 1.L.230.231; 1.L.230.236; 1.L.230.237; 1.L.230.238; 1.L.230.239; 1.L.230.154;
 1.L.230.157; 1.L.230.166; 1.L.230.169; 1.L.230.172; 1.L.230.175; 1.L.230.240;
 30 1.L.230.244; 1.L.231.228; 1.L.231.229; 1.L.231.230; 1.L.231.231; 1.L.231.236;

- 1.L.231.237; 1.L.231.238; 1.L.231.239; 1.L.231.154; 1.L.231.157; 1.L.231.166;
1.L.231.169; 1.L.231.172; 1.L.231.175; 1.L.231.240; 1.L.231.244; 1.L.236.228;
1.L.236.229; 1.L.236.230; 1.L.236.231; 1.L.236.236; 1.L.236.237; 1.L.236.238;
1.L.236.239; 1.L.236.154; 1.L.236.157; 1.L.236.166; 1.L.236.169; 1.L.236.172;
5 1.L.236.175; 1.L.236.240; 1.L.236.244; 1.L.237.228; 1.L.237.229; 1.L.237.230;
1.L.237.231; 1.L.237.236; 1.L.237.237; 1.L.237.238; 1.L.237.239; 1.L.237.154;
1.L.237.157; 1.L.237.166; 1.L.237.169; 1.L.237.172; 1.L.237.175; 1.L.237.240;
1.L.237.244; 1.L.238.228; 1.L.238.229; 1.L.238.230; 1.L.238.231; 1.L.238.236;
1.L.238.237; 1.L.238.238; 1.L.238.239; 1.L.238.154; 1.L.238.157; 1.L.238.166;
10 1.L.238.169; 1.L.238.172; 1.L.238.175; 1.L.238.240; 1.L.238.244; 1.L.239.228;
1.L.239.229; 1.L.239.230; 1.L.239.231; 1.L.239.236; 1.L.239.237; 1.L.239.238;
1.L.239.239; 1.L.239.154; 1.L.239.157; 1.L.239.166; 1.L.239.169; 1.L.239.172;
1.L.239.175; 1.L.239.240; 1.L.239.244; 1.L.154.228; 1.L.154.229; 1.L.154.230;
1.L.154.231; 1.L.154.236; 1.L.154.237; 1.L.154.238; 1.L.154.239; 1.L.154.154;
15 1.L.154.157; 1.L.154.166; 1.L.154.169; 1.L.154.172; 1.L.154.175; 1.L.154.240;
1.L.154.244; 1.L.157.228; 1.L.157.229; 1.L.157.230; 1.L.157.231; 1.L.157.236;
1.L.157.237; 1.L.157.238; 1.L.157.239; 1.L.157.154; 1.L.157.157; 1.L.157.166;
1.L.157.169; 1.L.157.172; 1.L.157.175; 1.L.157.240; 1.L.157.244; 1.L.166.228;
1.L.166.229; 1.L.166.230; 1.L.166.231; 1.L.166.236; 1.L.166.237; 1.L.166.238;
20 1.L.166.239; 1.L.166.154; 1.L.166.157; 1.L.166.166; 1.L.166.169; 1.L.166.172;
1.L.166.175; 1.L.166.240; 1.L.166.244; 1.L.169.228; 1.L.169.229; 1.L.169.230;
1.L.169.231; 1.L.169.236; 1.L.169.237; 1.L.169.238; 1.L.169.239; 1.L.169.154;
1.L.169.157; 1.L.169.166; 1.L.169.169; 1.L.169.172; 1.L.169.175; 1.L.169.240;
1.L.169.244; 1.L.172.228; 1.L.172.229; 1.L.172.230; 1.L.172.231; 1.L.172.236;
25 1.L.172.237; 1.L.172.238; 1.L.172.239; 1.L.172.154; 1.L.172.157; 1.L.172.166;
1.L.172.169; 1.L.172.172; 1.L.172.175; 1.L.172.240; 1.L.172.244; 1.L.175.228;
1.L.175.229; 1.L.175.230; 1.L.175.231; 1.L.175.236; 1.L.175.237; 1.L.175.238;
1.L.175.239; 1.L.175.154; 1.L.175.157; 1.L.175.166; 1.L.175.169; 1.L.175.172;
1.L.175.175; 1.L.175.240; 1.L.175.244; 1.L.240.228; 1.L.240.229; 1.L.240.230;
30 1.L.240.231; 1.L.240.236; 1.L.240.237; 1.L.240.238; 1.L.240.239; 1.L.240.154;

1.L.240.157; 1.L.240.166; 1.L.240.169; 1.L.240.172; 1.L.240.175; 1.L.240.240;
 1.L.240.244; 1.L.244.228; 1.L.244.229; 1.L.244.230; 1.L.244.231; 1.L.244.236;
 1.L.244.237; 1.L.244.238; 1.L.244.239; 1.L.244.154; 1.L.244.157; 1.L.244.166;
 1.L.244.169; 1.L.244.172; 1.L.244.175; 1.L.244.240; 1.L.244.244;

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Prodrugs of 1.O

1.O.228.228; 1.O.228.229; 1.O.228.230; 1.O.228.231; 1.O.228.236;
 1.O.228.237; 1.O.228.238; 1.O.228.239; 1.O.228.154; 1.O.228.157;
 1.O.228.166; 1.O.228.169; 1.O.228.172; 1.O.228.175; 1.O.228.240;
 10 1.O.228.244; 1.O.229.228; 1.O.229.229; 1.O.229.230; 1.O.229.231;
 1.O.229.236; 1.O.229.237; 1.O.229.238; 1.O.229.239; 1.O.229.154;
 1.O.229.157; 1.O.229.166; 1.O.229.169; 1.O.229.172; 1.O.229.175;
 1.O.229.240; 1.O.229.244; 1.O.230.228; 1.O.230.229; 1.O.230.230;
 1.O.230.231; 1.O.230.236; 1.O.230.237; 1.O.230.238; 1.O.230.239;
 15 1.O.230.154; 1.O.230.157; 1.O.230.166; 1.O.230.169; 1.O.230.172;
 1.O.230.175; 1.O.230.240; 1.O.230.244; 1.O.231.228; 1.O.231.229;
 1.O.231.230; 1.O.231.231; 1.O.231.236; 1.O.231.237; 1.O.231.238;
 1.O.231.239; 1.O.231.154; 1.O.231.157; 1.O.231.166; 1.O.231.169;
 1.O.231.172; 1.O.231.175; 1.O.231.240; 1.O.231.244; 1.O.236.228;
 20 1.O.236.229; 1.O.236.230; 1.O.236.231; 1.O.236.236; 1.O.236.237;
 1.O.236.238; 1.O.236.239; 1.O.236.154; 1.O.236.157; 1.O.236.166;
 1.O.236.169; 1.O.236.172; 1.O.236.175; 1.O.236.240; 1.O.236.244;
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 1.O.237.237; 1.O.237.238; 1.O.237.239; 1.O.237.154; 1.O.237.157;
 25 1.O.237.166; 1.O.237.169; 1.O.237.172; 1.O.237.175; 1.O.237.240;
 1.O.237.244; 1.O.238.228; 1.O.238.229; 1.O.238.230; 1.O.238.231;
 1.O.238.236; 1.O.238.237; 1.O.238.238; 1.O.238.239; 1.O.238.154;
 1.O.238.157; 1.O.238.166; 1.O.238.169; 1.O.238.172; 1.O.238.175;
 1.O.238.240; 1.O.238.244; 1.O.239.228; 1.O.239.229; 1.O.239.230;
 30 1.O.239.231; 1.O.239.236; 1.O.239.237; 1.O.239.238; 1.O.239.239;

1.O.239.154; 1.O.239.157; 1.O.239.166; 1.O.239.169; 1.O.239.172;
1.O.239.175; 1.O.239.240; 1.O.239.244; 1.O.154.228; 1.O.154.229;
1.O.154.230; 1.O.154.231; 1.O.154.236; 1.O.154.237; 1.O.154.238;
1.O.154.239; 1.O.154.154; 1.O.154.157; 1.O.154.166; 1.O.154.169;
5 1.O.154.172; 1.O.154.175; 1.O.154.240; 1.O.154.244; 1.O.157.228;
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1.O.157.238; 1.O.157.239; 1.O.157.154; 1.O.157.157; 1.O.157.166;
1.O.157.169; 1.O.157.172; 1.O.157.175; 1.O.157.240; 1.O.157.244;
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10 1.O.166.237; 1.O.166.238; 1.O.166.239; 1.O.166.154; 1.O.166.157;
1.O.166.166; 1.O.166.169; 1.O.166.172; 1.O.166.175; 1.O.166.240;
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1.O.169.236; 1.O.169.237; 1.O.169.238; 1.O.169.239; 1.O.169.154;
1.O.169.157; 1.O.169.166; 1.O.169.169; 1.O.169.172; 1.O.169.175;
15 1.O.169.240; 1.O.169.244; 1.O.172.228; 1.O.172.229; 1.O.172.230;
1.O.172.231; 1.O.172.236; 1.O.172.237; 1.O.172.238; 1.O.172.239;
1.O.172.154; 1.O.172.157; 1.O.172.166; 1.O.172.169; 1.O.172.172;
1.O.172.175; 1.O.172.240; 1.O.172.244; 1.O.175.228; 1.O.175.229;
1.O.175.230; 1.O.175.231; 1.O.175.236; 1.O.175.237; 1.O.175.238;
20 1.O.175.239; 1.O.175.154; 1.O.175.157; 1.O.175.166; 1.O.175.169;
1.O.175.172; 1.O.175.175; 1.O.175.240; 1.O.175.244; 1.O.240.228;
1.O.240.229; 1.O.240.230; 1.O.240.231; 1.O.240.236; 1.O.240.237;
1.O.240.238; 1.O.240.239; 1.O.240.154; 1.O.240.157; 1.O.240.166;
1.O.240.169; 1.O.240.172; 1.O.240.175; 1.O.240.240; 1.O.240.244;
25 1.O.244.228; 1.O.244.229; 1.O.244.230; 1.O.244.231; 1.O.244.236;
1.O.244.237; 1.O.244.238; 1.O.244.239; 1.O.244.154; 1.O.244.157;
1.O.244.166; 1.O.244.169; 1.O.244.172; 1.O.244.175; 1.O.244.240;
1.O.244.244;

30 Prodrugs of 1.P

1.P.228.228; 1.P.228.229; 1.P.228.230; 1.P.228.231; 1.P.228.236;
1.P.228.237; 1.P.228.238; 1.P.228.239; 1.P.228.154; 1.P.228.157; 1.P.228.166;
1.P.228.169; 1.P.228.172; 1.P.228.175; 1.P.228.240; 1.P.228.244; 1.P.229.228;
1.P.229.229; 1.P.229.230; 1.P.229.231; 1.P.229.236; 1.P.229.237; 1.P.229.238;
5 1.P.229.239; 1.P.229.154; 1.P.229.157; 1.P.229.166; 1.P.229.169; 1.P.229.172;
1.P.229.175; 1.P.229.240; 1.P.229.244; 1.P.230.228; 1.P.230.229; 1.P.230.230;
1.P.230.231; 1.P.230.236; 1.P.230.237; 1.P.230.238; 1.P.230.239; 1.P.230.154;
1.P.230.157; 1.P.230.166; 1.P.230.169; 1.P.230.172; 1.P.230.175; 1.P.230.240;
1.P.230.244; 1.P.231.228; 1.P.231.229; 1.P.231.230; 1.P.231.231; 1.P.231.236;
10 1.P.231.237; 1.P.231.238; 1.P.231.239; 1.P.231.154; 1.P.231.157; 1.P.231.166;
1.P.231.169; 1.P.231.172; 1.P.231.175; 1.P.231.240; 1.P.231.244; 1.P.236.228;
1.P.236.229; 1.P.236.230; 1.P.236.231; 1.P.236.236; 1.P.236.237; 1.P.236.238;
1.P.236.239; 1.P.236.154; 1.P.236.157; 1.P.236.166; 1.P.236.169; 1.P.236.172;
1.P.236.175; 1.P.236.240; 1.P.236.244; 1.P.237.228; 1.P.237.229; 1.P.237.230;
15 1.P.237.231; 1.P.237.236; 1.P.237.237; 1.P.237.238; 1.P.237.239; 1.P.237.154;
1.P.237.157; 1.P.237.166; 1.P.237.169; 1.P.237.172; 1.P.237.175; 1.P.237.240;
1.P.237.244; 1.P.238.228; 1.P.238.229; 1.P.238.230; 1.P.238.231; 1.P.238.236;
1.P.238.237; 1.P.238.238; 1.P.238.239; 1.P.238.154; 1.P.238.157; 1.P.238.166;
1.P.238.169; 1.P.238.172; 1.P.238.175; 1.P.238.240; 1.P.238.244; 1.P.239.228;
20 1.P.239.229; 1.P.239.230; 1.P.239.231; 1.P.239.236; 1.P.239.237; 1.P.239.238;
1.P.239.239; 1.P.239.154; 1.P.239.157; 1.P.239.166; 1.P.239.169; 1.P.239.172;
1.P.239.175; 1.P.239.240; 1.P.239.244; 1.P.154.228; 1.P.154.229; 1.P.154.230;
1.P.154.231; 1.P.154.236; 1.P.154.237; 1.P.154.238; 1.P.154.239; 1.P.154.154;
1.P.154.157; 1.P.154.166; 1.P.154.169; 1.P.154.172; 1.P.154.175; 1.P.154.240;
25 1.P.154.244; 1.P.157.228; 1.P.157.229; 1.P.157.230; 1.P.157.231; 1.P.157.236;
1.P.157.237; 1.P.157.238; 1.P.157.239; 1.P.157.154; 1.P.157.157; 1.P.157.166;
1.P.157.169; 1.P.157.172; 1.P.157.175; 1.P.157.240; 1.P.157.244; 1.P.166.228;
1.P.166.229; 1.P.166.230; 1.P.166.231; 1.P.166.236; 1.P.166.237; 1.P.166.238;
1.P.166.239; 1.P.166.154; 1.P.166.157; 1.P.166.166; 1.P.166.169; 1.P.166.172;
30 1.P.166.175; 1.P.166.240; 1.P.166.244; 1.P.169.228; 1.P.169.229; 1.P.169.230;

1.P.169.231; 1.P.169.236; 1.P.169.237; 1.P.169.238; 1.P.169.239; 1.P.169.154;
1.P.169.157; 1.P.169.166; 1.P.169.169; 1.P.169.172; 1.P.169.175; 1.P.169.240;
1.P.169.244; 1.P.172.228; 1.P.172.229; 1.P.172.230; 1.P.172.231; 1.P.172.236;
1.P.172.237; 1.P.172.238; 1.P.172.239; 1.P.172.154; 1.P.172.157; 1.P.172.166;
5 1.P.172.169; 1.P.172.172; 1.P.172.175; 1.P.172.240; 1.P.172.244; 1.P.175.228;
1.P.175.229; 1.P.175.230; 1.P.175.231; 1.P.175.236; 1.P.175.237; 1.P.175.238;
1.P.175.239; 1.P.175.154; 1.P.175.157; 1.P.175.166; 1.P.175.169; 1.P.175.172;
1.P.175.175; 1.P.175.240; 1.P.175.244; 1.P.240.228; 1.P.240.229; 1.P.240.230;
1.P.240.231; 1.P.240.236; 1.P.240.237; 1.P.240.238; 1.P.240.239; 1.P.240.154;
10 1.P.240.157; 1.P.240.166; 1.P.240.169; 1.P.240.172; 1.P.240.175; 1.P.240.240;
1.P.240.244; 1.P.244.228; 1.P.244.229; 1.P.244.230; 1.P.244.231; 1.P.244.236;
1.P.244.237; 1.P.244.238; 1.P.244.239; 1.P.244.154; 1.P.244.157; 1.P.244.166;
1.P.244.169; 1.P.244.172; 1.P.244.175; 1.P.244.240; 1.P.244.244;

15 Prodrugs of 1.U

1.U.228.228; 1.U.228.229; 1.U.228.230; 1.U.228.231; 1.U.228.236;
1.U.228.237; 1.U.228.238; 1.U.228.239; 1.U.228.154; 1.U.228.157;
1.U.228.166; 1.U.228.169; 1.U.228.172; 1.U.228.175; 1.U.228.240;
1.U.228.244; 1.U.229.228; 1.U.229.229; 1.U.229.230; 1.U.229.231;
20 1.U.229.236; 1.U.229.237; 1.U.229.238; 1.U.229.239; 1.U.229.154;
1.U.229.157; 1.U.229.166; 1.U.229.169; 1.U.229.172; 1.U.229.175;
1.U.229.240; 1.U.229.244; 1.U.230.228; 1.U.230.229; 1.U.230.230;
1.U.230.231; 1.U.230.236; 1.U.230.237; 1.U.230.238; 1.U.230.239;
1.U.230.154; 1.U.230.157; 1.U.230.166; 1.U.230.169; 1.U.230.172;
25 1.U.230.175; 1.U.230.240; 1.U.230.244; 1.U.231.228; 1.U.231.229;
1.U.231.230; 1.U.231.231; 1.U.231.236; 1.U.231.237; 1.U.231.238;
1.U.231.239; 1.U.231.154; 1.U.231.157; 1.U.231.166; 1.U.231.169;
1.U.231.172; 1.U.231.175; 1.U.231.240; 1.U.231.244; 1.U.236.228;
1.U.236.229; 1.U.236.230; 1.U.236.231; 1.U.236.236; 1.U.236.237;
30 1.U.236.238; 1.U.236.239; 1.U.236.154; 1.U.236.157; 1.U.236.166;

1.U.236.169; 1.U.236.172; 1.U.236.175; 1.U.236.240; 1.U.236.244;
1.U.237.228; 1.U.237.229; 1.U.237.230; 1.U.237.231; 1.U.237.236;
1.U.237.237; 1.U.237.238; 1.U.237.239; 1.U.237.154; 1.U.237.157;
1.U.237.166; 1.U.237.169; 1.U.237.172; 1.U.237.175; 1.U.237.240;
5 1.U.237.244; 1.U.238.228; 1.U.238.229; 1.U.238.230; 1.U.238.231;
1.U.238.236; 1.U.238.237; 1.U.238.238; 1.U.238.239; 1.U.238.154;
1.U.238.157; 1.U.238.166; 1.U.238.169; 1.U.238.172; 1.U.238.175;
1.U.238.240; 1.U.238.244; 1.U.239.228; 1.U.239.229; 1.U.239.230;
1.U.239.231; 1.U.239.236; 1.U.239.237; 1.U.239.238; 1.U.239.239;
10 1.U.239.154; 1.U.239.157; 1.U.239.166; 1.U.239.169; 1.U.239.172;
1.U.239.175; 1.U.239.240; 1.U.239.244; 1.U.154.228; 1.U.154.229;
1.U.154.230; 1.U.154.231; 1.U.154.236; 1.U.154.237; 1.U.154.238;
1.U.154.239; 1.U.154.154; 1.U.154.157; 1.U.154.166; 1.U.154.169;
1.U.154.172; 1.U.154.175; 1.U.154.240; 1.U.154.244; 1.U.157.228;
15 1.U.157.229; 1.U.157.230; 1.U.157.231; 1.U.157.236; 1.U.157.237;
1.U.157.238; 1.U.157.239; 1.U.157.154; 1.U.157.157; 1.U.157.166;
1.U.157.169; 1.U.157.172; 1.U.157.175; 1.U.157.240; 1.U.157.244;
1.U.166.228; 1.U.166.229; 1.U.166.230; 1.U.166.231; 1.U.166.236;
1.U.166.237; 1.U.166.238; 1.U.166.239; 1.U.166.154; 1.U.166.157;
20 1.U.166.166; 1.U.166.169; 1.U.166.172; 1.U.166.175; 1.U.166.240;
1.U.166.244; 1.U.169.228; 1.U.169.229; 1.U.169.230; 1.U.169.231;
1.U.169.236; 1.U.169.237; 1.U.169.238; 1.U.169.239; 1.U.169.154;
1.U.169.157; 1.U.169.166; 1.U.169.169; 1.U.169.172; 1.U.169.175;
1.U.169.240; 1.U.169.244; 1.U.172.228; 1.U.172.229; 1.U.172.230;
25 1.U.172.231; 1.U.172.236; 1.U.172.237; 1.U.172.238; 1.U.172.239;
1.U.172.154; 1.U.172.157; 1.U.172.166; 1.U.172.169; 1.U.172.172;
1.U.172.175; 1.U.172.240; 1.U.172.244; 1.U.175.228; 1.U.175.229;
1.U.175.230; 1.U.175.231; 1.U.175.236; 1.U.175.237; 1.U.175.238;
1.U.175.239; 1.U.175.154; 1.U.175.157; 1.U.175.166; 1.U.175.169;
30 1.U.175.172; 1.U.175.175; 1.U.175.240; 1.U.175.244; 1.U.240.228;

1.U.240.229; 1.U.240.230; 1.U.240.231; 1.U.240.236; 1.U.240.237;
1.U.240.238; 1.U.240.239; 1.U.240.154; 1.U.240.157; 1.U.240.166;
1.U.240.169; 1.U.240.172; 1.U.240.175; 1.U.240.240; 1.U.240.244;
1.U.244.228; 1.U.244.229; 1.U.244.230; 1.U.244.231; 1.U.244.236;
5 1.U.244.237; 1.U.244.238; 1.U.244.239; 1.U.244.154; 1.U.244.157;
1.U.244.166; 1.U.244.169; 1.U.244.172; 1.U.244.175; 1.U.244.240;
1.U.244.244;

Prodrugs of 1.W

10 1.W.228.228; 1.W.228.229; 1.W.228.230; 1.W.228.231; 1.W.228.236;
1.W.228.237; 1.W.228.238; 1.W.228.239; 1.W.228.154; 1.W.228.157;
1.W.228.166; 1.W.228.169; 1.W.228.172; 1.W.228.175; 1.W.228.240;
1.W.228.244; 1.W.229.228; 1.W.229.229; 1.W.229.230; 1.W.229.231;
1.W.229.236; 1.W.229.237; 1.W.229.238; 1.W.229.239; 1.W.229.154;
15 1.W.229.157; 1.W.229.166; 1.W.229.169; 1.W.229.172; 1.W.229.175;
1.W.229.240; 1.W.229.244; 1.W.230.228; 1.W.230.229; 1.W.230.230;
1.W.230.231; 1.W.230.236; 1.W.230.237; 1.W.230.238; 1.W.230.239;
1.W.230.154; 1.W.230.157; 1.W.230.166; 1.W.230.169; 1.W.230.172;
1.W.230.175; 1.W.230.240; 1.W.230.244; 1.W.231.228; 1.W.231.229;
20 1.W.231.230; 1.W.231.231; 1.W.231.236; 1.W.231.237; 1.W.231.238;
1.W.231.239; 1.W.231.154; 1.W.231.157; 1.W.231.166; 1.W.231.169;
1.W.231.172; 1.W.231.175; 1.W.231.240; 1.W.231.244; 1.W.236.228;
1.W.236.229; 1.W.236.230; 1.W.236.231; 1.W.236.236; 1.W.236.237;
1.W.236.238; 1.W.236.239; 1.W.236.154; 1.W.236.157; 1.W.236.166;
25 1.W.236.169; 1.W.236.172; 1.W.236.175; 1.W.236.240; 1.W.236.244;
1.W.237.228; 1.W.237.229; 1.W.237.230; 1.W.237.231; 1.W.237.236;
1.W.237.237; 1.W.237.238; 1.W.237.239; 1.W.237.154; 1.W.237.157;
1.W.237.166; 1.W.237.169; 1.W.237.172; 1.W.237.175; 1.W.237.240;
1.W.237.244; 1.W.238.228; 1.W.238.229; 1.W.238.230; 1.W.238.231;
30 1.W.238.236; 1.W.238.237; 1.W.238.238; 1.W.238.239; 1.W.238.154;

1.W.238.157; 1.W.238.166; 1.W.238.169; 1.W.238.172; 1.W.238.175;
1.W.238.240; 1.W.238.244; 1.W.239.228; 1.W.239.229; 1.W.239.230;
1.W.239.231; 1.W.239.236; 1.W.239.237; 1.W.239.238; 1.W.239.239;
1.W.239.154; 1.W.239.157; 1.W.239.166; 1.W.239.169; 1.W.239.172;
5 1.W.239.175; 1.W.239.240; 1.W.239.244; 1.W.154.228; 1.W.154.229;
1.W.154.230; 1.W.154.231; 1.W.154.236; 1.W.154.237; 1.W.154.238;
1.W.154.239; 1.W.154.154; 1.W.154.157; 1.W.154.166; 1.W.154.169;
1.W.154.172; 1.W.154.175; 1.W.154.240; 1.W.154.244; 1.W.157.228;
1.W.157.229; 1.W.157.230; 1.W.157.231; 1.W.157.236; 1.W.157.237;
10 1.W.157.238; 1.W.157.239; 1.W.157.154; 1.W.157.157; 1.W.157.166;
1.W.157.169; 1.W.157.172; 1.W.157.175; 1.W.157.240; 1.W.157.244;
1.W.166.228; 1.W.166.229; 1.W.166.230; 1.W.166.231; 1.W.166.236;
1.W.166.237; 1.W.166.238; 1.W.166.239; 1.W.166.154; 1.W.166.157;
1.W.166.166; 1.W.166.169; 1.W.166.172; 1.W.166.175; 1.W.166.240;
15 1.W.166.244; 1.W.169.228; 1.W.169.229; 1.W.169.230; 1.W.169.231;
1.W.169.236; 1.W.169.237; 1.W.169.238; 1.W.169.239; 1.W.169.154;
1.W.169.157; 1.W.169.166; 1.W.169.169; 1.W.169.172; 1.W.169.175;
1.W.169.240; 1.W.169.244; 1.W.172.228; 1.W.172.229; 1.W.172.230;
1.W.172.231; 1.W.172.236; 1.W.172.237; 1.W.172.238; 1.W.172.239;
20 1.W.172.154; 1.W.172.157; 1.W.172.166; 1.W.172.169; 1.W.172.172;
1.W.172.175; 1.W.172.240; 1.W.172.244; 1.W.175.228; 1.W.175.229;
1.W.175.230; 1.W.175.231; 1.W.175.236; 1.W.175.237; 1.W.175.238;
1.W.175.239; 1.W.175.154; 1.W.175.157; 1.W.175.166; 1.W.175.169;
1.W.175.172; 1.W.175.175; 1.W.175.240; 1.W.175.244; 1.W.240.228;
25 1.W.240.229; 1.W.240.230; 1.W.240.231; 1.W.240.236; 1.W.240.237;
1.W.240.238; 1.W.240.239; 1.W.240.154; 1.W.240.157; 1.W.240.166;
1.W.240.169; 1.W.240.172; 1.W.240.175; 1.W.240.240; 1.W.240.244;
1.W.244.228; 1.W.244.229; 1.W.244.230; 1.W.244.231; 1.W.244.236;
1.W.244.237; 1.W.244.238; 1.W.244.239; 1.W.244.154; 1.W.244.157;

1.W.244.166; 1.W.244.169; 1.W.244.172; 1.W.244.175; 1.W.244.240;
1.W.244.244;

Prodrugs of 1.Y

- 5 1.Y.228.228; 1.Y.228.229; 1.Y.228.230; 1.Y.228.231; 1.Y.228.236;
1.Y.228.237; 1.Y.228.238; 1.Y.228.239; 1.Y.228.154; 1.Y.228.157; 1.Y.228.166;
1.Y.228.169; 1.Y.228.172; 1.Y.228.175; 1.Y.228.240; 1.Y.228.244; 1.Y.229.228;
1.Y.229.229; 1.Y.229.230; 1.Y.229.231; 1.Y.229.236; 1.Y.229.237; 1.Y.229.238;
1.Y.229.239; 1.Y.229.154; 1.Y.229.157; 1.Y.229.166; 1.Y.229.169; 1.Y.229.172;
10 1.Y.229.175; 1.Y.229.240; 1.Y.229.244; 1.Y.230.228; 1.Y.230.229; 1.Y.230.230;
1.Y.230.231; 1.Y.230.236; 1.Y.230.237; 1.Y.230.238; 1.Y.230.239; 1.Y.230.154;
1.Y.230.157; 1.Y.230.166; 1.Y.230.169; 1.Y.230.172; 1.Y.230.175; 1.Y.230.240;
1.Y.230.244; 1.Y.231.228; 1.Y.231.229; 1.Y.231.230; 1.Y.231.231; 1.Y.231.236;
1.Y.231.237; 1.Y.231.238; 1.Y.231.239; 1.Y.231.154; 1.Y.231.157; 1.Y.231.166;
15 1.Y.231.169; 1.Y.231.172; 1.Y.231.175; 1.Y.231.240; 1.Y.231.244; 1.Y.236.228;
1.Y.236.229; 1.Y.236.230; 1.Y.236.231; 1.Y.236.236; 1.Y.236.237; 1.Y.236.238;
1.Y.236.239; 1.Y.236.154; 1.Y.236.157; 1.Y.236.166; 1.Y.236.169; 1.Y.236.172;
1.Y.236.175; 1.Y.236.240; 1.Y.236.244; 1.Y.237.228; 1.Y.237.229; 1.Y.237.230;
1.Y.237.231; 1.Y.237.236; 1.Y.237.237; 1.Y.237.238; 1.Y.237.239; 1.Y.237.154;
20 1.Y.237.157; 1.Y.237.166; 1.Y.237.169; 1.Y.237.172; 1.Y.237.175; 1.Y.237.240;
1.Y.237.244; 1.Y.238.228; 1.Y.238.229; 1.Y.238.230; 1.Y.238.231; 1.Y.238.236;
1.Y.238.237; 1.Y.238.238; 1.Y.238.239; 1.Y.238.154; 1.Y.238.157; 1.Y.238.166;
1.Y.238.169; 1.Y.238.172; 1.Y.238.175; 1.Y.238.240; 1.Y.238.244; 1.Y.239.228;
1.Y.239.229; 1.Y.239.230; 1.Y.239.231; 1.Y.239.236; 1.Y.239.237; 1.Y.239.238;
25 1.Y.239.239; 1.Y.239.154; 1.Y.239.157; 1.Y.239.166; 1.Y.239.169; 1.Y.239.172;
1.Y.239.175; 1.Y.239.240; 1.Y.239.244; 1.Y.154.228; 1.Y.154.229; 1.Y.154.230;
1.Y.154.231; 1.Y.154.236; 1.Y.154.237; 1.Y.154.238; 1.Y.154.239; 1.Y.154.154;
1.Y.154.157; 1.Y.154.166; 1.Y.154.169; 1.Y.154.172; 1.Y.154.175; 1.Y.154.240;
1.Y.154.244; 1.Y.157.228; 1.Y.157.229; 1.Y.157.230; 1.Y.157.231; 1.Y.157.236;
30 1.Y.157.237; 1.Y.157.238; 1.Y.157.239; 1.Y.157.154; 1.Y.157.157; 1.Y.157.166;

1.Y.157.169; 1.Y.157.172; 1.Y.157.175; 1.Y.157.240; 1.Y.157.244; 1.Y.166.228;
 1.Y.166.229; 1.Y.166.230; 1.Y.166.231; 1.Y.166.236; 1.Y.166.237; 1.Y.166.238;
 1.Y.166.239; 1.Y.166.154; 1.Y.166.157; 1.Y.166.166; 1.Y.166.169; 1.Y.166.172;
 1.Y.166.175; 1.Y.166.240; 1.Y.166.244; 1.Y.169.228; 1.Y.169.229; 1.Y.169.230;
 5 1.Y.169.231; 1.Y.169.236; 1.Y.169.237; 1.Y.169.238; 1.Y.169.239; 1.Y.169.154;
 1.Y.169.157; 1.Y.169.166; 1.Y.169.169; 1.Y.169.172; 1.Y.169.175; 1.Y.169.240;
 1.Y.169.244; 1.Y.172.228; 1.Y.172.229; 1.Y.172.230; 1.Y.172.231; 1.Y.172.236;
 1.Y.172.237; 1.Y.172.238; 1.Y.172.239; 1.Y.172.154; 1.Y.172.157; 1.Y.172.166;
 1.Y.172.169; 1.Y.172.172; 1.Y.172.175; 1.Y.172.240; 1.Y.172.244; 1.Y.175.228;
 10 1.Y.175.229; 1.Y.175.230; 1.Y.175.231; 1.Y.175.236; 1.Y.175.237; 1.Y.175.238;
 1.Y.175.239; 1.Y.175.154; 1.Y.175.157; 1.Y.175.166; 1.Y.175.169; 1.Y.175.172;
 1.Y.175.175; 1.Y.175.240; 1.Y.175.244; 1.Y.240.228; 1.Y.240.229; 1.Y.240.230;
 1.Y.240.231; 1.Y.240.236; 1.Y.240.237; 1.Y.240.238; 1.Y.240.239; 1.Y.240.154;
 1.Y.240.157; 1.Y.240.166; 1.Y.240.169; 1.Y.240.172; 1.Y.240.175; 1.Y.240.240;
 15 1.Y.240.244; 1.Y.244.228; 1.Y.244.229; 1.Y.244.230; 1.Y.244.231; 1.Y.244.236;
 1.Y.244.237; 1.Y.244.238; 1.Y.244.239; 1.Y.244.154; 1.Y.244.157; 1.Y.244.166;
 1.Y.244.169; 1.Y.244.172; 1.Y.244.175; 1.Y.244.240; 1.Y.244.244;

Prodrugs of 2.B

20 2.B.228.228; 2.B.228.229; 2.B.228.230; 2.B.228.231; 2.B.228.236;
 2.B.228.237; 2.B.228.238; 2.B.228.239; 2.B.228.154; 2.B.228.157; 2.B.228.166;
 2.B.228.169; 2.B.228.172; 2.B.228.175; 2.B.228.240; 2.B.228.244; 2.B.229.228;
 2.B.229.229; 2.B.229.230; 2.B.229.231; 2.B.229.236; 2.B.229.237; 2.B.229.238;
 2.B.229.239; 2.B.229.154; 2.B.229.157; 2.B.229.166; 2.B.229.169; 2.B.229.172;
 25 2.B.229.175; 2.B.229.240; 2.B.229.244; 2.B.230.228; 2.B.230.229; 2.B.230.230;
 2.B.230.231; 2.B.230.236; 2.B.230.237; 2.B.230.238; 2.B.230.239; 2.B.230.154;
 2.B.230.157; 2.B.230.166; 2.B.230.169; 2.B.230.172; 2.B.230.175; 2.B.230.240;
 2.B.230.244; 2.B.231.228; 2.B.231.229; 2.B.231.230; 2.B.231.231; 2.B.231.236;
 2.B.231.237; 2.B.231.238; 2.B.231.239; 2.B.231.154; 2.B.231.157; 2.B.231.166;
 30 2.B.231.169; 2.B.231.172; 2.B.231.175; 2.B.231.240; 2.B.231.244; 2.B.236.228;

2.B.236.229; 2.B.236.230; 2.B.236.231; 2.B.236.236; 2.B.236.237; 2.B.236.238;
2.B.236.239; 2.B.236.154; 2.B.236.157; 2.B.236.166; 2.B.236.169; 2.B.236.172;
2.B.236.175; 2.B.236.240; 2.B.236.244; 2.B.237.228; 2.B.237.229; 2.B.237.230;
2.B.237.231; 2.B.237.236; 2.B.237.237; 2.B.237.238; 2.B.237.239; 2.B.237.154;
5 2.B.237.157; 2.B.237.166; 2.B.237.169; 2.B.237.172; 2.B.237.175; 2.B.237.240;
2.B.237.244; 2.B.238.228; 2.B.238.229; 2.B.238.230; 2.B.238.231; 2.B.238.236;
2.B.238.237; 2.B.238.238; 2.B.238.239; 2.B.238.154; 2.B.238.157; 2.B.238.166;
2.B.238.169; 2.B.238.172; 2.B.238.175; 2.B.238.240; 2.B.238.244; 2.B.239.228;
2.B.239.229; 2.B.239.230; 2.B.239.231; 2.B.239.236; 2.B.239.237; 2.B.239.238;
10 2.B.239.239; 2.B.239.154; 2.B.239.157; 2.B.239.166; 2.B.239.169; 2.B.239.172;
2.B.239.175; 2.B.239.240; 2.B.239.244; 2.B.154.228; 2.B.154.229; 2.B.154.230;
2.B.154.231; 2.B.154.236; 2.B.154.237; 2.B.154.238; 2.B.154.239; 2.B.154.154;
2.B.154.157; 2.B.154.166; 2.B.154.169; 2.B.154.172; 2.B.154.175; 2.B.154.240;
2.B.154.244; 2.B.157.228; 2.B.157.229; 2.B.157.230; 2.B.157.231; 2.B.157.236;
15 2.B.157.237; 2.B.157.238; 2.B.157.239; 2.B.157.154; 2.B.157.157; 2.B.157.166;
2.B.157.169; 2.B.157.172; 2.B.157.175; 2.B.157.240; 2.B.157.244; 2.B.166.228;
2.B.166.229; 2.B.166.230; 2.B.166.231; 2.B.166.236; 2.B.166.237; 2.B.166.238;
2.B.166.239; 2.B.166.154; 2.B.166.157; 2.B.166.166; 2.B.166.169; 2.B.166.172;
2.B.166.175; 2.B.166.240; 2.B.166.244; 2.B.169.228; 2.B.169.229; 2.B.169.230;
20 2.B.169.231; 2.B.169.236; 2.B.169.237; 2.B.169.238; 2.B.169.239; 2.B.169.154;
2.B.169.157; 2.B.169.166; 2.B.169.169; 2.B.169.172; 2.B.169.175; 2.B.169.240;
2.B.169.244; 2.B.172.228; 2.B.172.229; 2.B.172.230; 2.B.172.231; 2.B.172.236;
2.B.172.237; 2.B.172.238; 2.B.172.239; 2.B.172.154; 2.B.172.157; 2.B.172.166;
2.B.172.169; 2.B.172.172; 2.B.172.175; 2.B.172.240; 2.B.172.244; 2.B.175.228;
25 2.B.175.229; 2.B.175.230; 2.B.175.231; 2.B.175.236; 2.B.175.237; 2.B.175.238;
2.B.175.239; 2.B.175.154; 2.B.175.157; 2.B.175.166; 2.B.175.169; 2.B.175.172;
2.B.175.175; 2.B.175.240; 2.B.175.244; 2.B.240.228; 2.B.240.229; 2.B.240.230;
2.B.240.231; 2.B.240.236; 2.B.240.237; 2.B.240.238; 2.B.240.239; 2.B.240.154;
2.B.240.157; 2.B.240.166; 2.B.240.169; 2.B.240.172; 2.B.240.175; 2.B.240.240;
30 2.B.240.244; 2.B.244.228; 2.B.244.229; 2.B.244.230; 2.B.244.231; 2.B.244.236;

2.B.244.237; 2.B.244.238; 2.B.244.239; 2.B.244.154; 2.B.244.157; 2.B.244.166;
2.B.244.169; 2.B.244.172; 2.B.244.175; 2.B.244.240; 2.B.244.244;

Prodrugs of 2.D

- 5 2.D.228.228; 2.D.228.229; 2.D.228.230; 2.D.228.231; 2.D.228.236;
2.D.228.237; 2.D.228.238; 2.D.228.239; 2.D.228.154; 2.D.228.157;
2.D.228.166; 2.D.228.169; 2.D.228.172; 2.D.228.175; 2.D.228.240;
2.D.228.244; 2.D.229.228; 2.D.229.229; 2.D.229.230; 2.D.229.231;
2.D.229.236; 2.D.229.237; 2.D.229.238; 2.D.229.239; 2.D.229.154;
10 2.D.229.157; 2.D.229.166; 2.D.229.169; 2.D.229.172; 2.D.229.175;
2.D.229.240; 2.D.229.244; 2.D.230.228; 2.D.230.229; 2.D.230.230;
2.D.230.231; 2.D.230.236; 2.D.230.237; 2.D.230.238; 2.D.230.239;
2.D.230.154; 2.D.230.157; 2.D.230.166; 2.D.230.169; 2.D.230.172;
2.D.230.175; 2.D.230.240; 2.D.230.244; 2.D.231.228; 2.D.231.229;
15 2.D.231.230; 2.D.231.231; 2.D.231.236; 2.D.231.237; 2.D.231.238;
2.D.231.239; 2.D.231.154; 2.D.231.157; 2.D.231.166; 2.D.231.169;
2.D.231.172; 2.D.231.175; 2.D.231.240; 2.D.231.244; 2.D.236.228;
2.D.236.229; 2.D.236.230; 2.D.236.231; 2.D.236.236; 2.D.236.237;
2.D.236.238; 2.D.236.239; 2.D.236.154; 2.D.236.157; 2.D.236.166;
20 2.D.236.169; 2.D.236.172; 2.D.236.175; 2.D.236.240; 2.D.236.244;
2.D.237.228; 2.D.237.229; 2.D.237.230; 2.D.237.231; 2.D.237.236;
2.D.237.237; 2.D.237.238; 2.D.237.239; 2.D.237.154; 2.D.237.157;
2.D.237.166; 2.D.237.169; 2.D.237.172; 2.D.237.175; 2.D.237.240;
2.D.237.244; 2.D.238.228; 2.D.238.229; 2.D.238.230; 2.D.238.231;
25 2.D.238.236; 2.D.238.237; 2.D.238.238; 2.D.238.239; 2.D.238.154;
2.D.238.157; 2.D.238.166; 2.D.238.169; 2.D.238.172; 2.D.238.175;
2.D.238.240; 2.D.238.244; 2.D.239.228; 2.D.239.229; 2.D.239.230;
2.D.239.231; 2.D.239.236; 2.D.239.237; 2.D.239.238; 2.D.239.239;
2.D.239.154; 2.D.239.157; 2.D.239.166; 2.D.239.169; 2.D.239.172;
30 2.D.239.175; 2.D.239.240; 2.D.239.244; 2.D.154.228; 2.D.154.229;

2.D.154.230; 2.D.154.231; 2.D.154.236; 2.D.154.237; 2.D.154.238;
2.D.154.239; 2.D.154.154; 2.D.154.157; 2.D.154.166; 2.D.154.169;
2.D.154.172; 2.D.154.175; 2.D.154.240; 2.D.154.244; 2.D.157.228;
2.D.157.229; 2.D.157.230; 2.D.157.231; 2.D.157.236; 2.D.157.237;
5 2.D.157.238; 2.D.157.239; 2.D.157.154; 2.D.157.157; 2.D.157.166;
2.D.157.169; 2.D.157.172; 2.D.157.175; 2.D.157.240; 2.D.157.244;
2.D.166.228; 2.D.166.229; 2.D.166.230; 2.D.166.231; 2.D.166.236;
2.D.166.237; 2.D.166.238; 2.D.166.239; 2.D.166.154; 2.D.166.157;
2.D.166.166; 2.D.166.169; 2.D.166.172; 2.D.166.175; 2.D.166.240;
10 2.D.166.244; 2.D.169.228; 2.D.169.229; 2.D.169.230; 2.D.169.231;
2.D.169.236; 2.D.169.237; 2.D.169.238; 2.D.169.239; 2.D.169.154;
2.D.169.157; 2.D.169.166; 2.D.169.169; 2.D.169.172; 2.D.169.175;
2.D.169.240; 2.D.169.244; 2.D.172.228; 2.D.172.229; 2.D.172.230;
2.D.172.231; 2.D.172.236; 2.D.172.237; 2.D.172.238; 2.D.172.239;
15 2.D.172.154; 2.D.172.157; 2.D.172.166; 2.D.172.169; 2.D.172.172;
2.D.172.175; 2.D.172.240; 2.D.172.244; 2.D.175.228; 2.D.175.229;
2.D.175.230; 2.D.175.231; 2.D.175.236; 2.D.175.237; 2.D.175.238;
2.D.175.239; 2.D.175.154; 2.D.175.157; 2.D.175.166; 2.D.175.169;
2.D.175.172; 2.D.175.175; 2.D.175.240; 2.D.175.244; 2.D.240.228;
20 2.D.240.229; 2.D.240.230; 2.D.240.231; 2.D.240.236; 2.D.240.237;
2.D.240.238; 2.D.240.239; 2.D.240.154; 2.D.240.157; 2.D.240.166;
2.D.240.169; 2.D.240.172; 2.D.240.175; 2.D.240.240; 2.D.240.244;
2.D.244.228; 2.D.244.229; 2.D.244.230; 2.D.244.231; 2.D.244.236;
2.D.244.237; 2.D.244.238; 2.D.244.239; 2.D.244.154; 2.D.244.157;
25 2.D.244.166; 2.D.244.169; 2.D.244.172; 2.D.244.175; 2.D.244.240;
2.D.244.244;

Prodrugs of 2.E

2.E.228.228; 2.E.228.229; 2.E.228.230; 2.E.228.231; 2.E.228.236;
30 2.E.228.237; 2.E.228.238; 2.E.228.239; 2.E.228.154; 2.E.228.157; 2.E.228.166;

2.E.228.169; 2.E.228.172; 2.E.228.175; 2.E.228.240; 2.E.228.244; 2.E.229.228;
2.E.229.229; 2.E.229.230; 2.E.229.231; 2.E.229.236; 2.E.229.237; 2.E.229.238;
2.E.229.239; 2.E.229.154; 2.E.229.157; 2.E.229.166; 2.E.229.169; 2.E.229.172;
2.E.229.175; 2.E.229.240; 2.E.229.244; 2.E.230.228; 2.E.230.229; 2.E.230.230;
5 2.E.230.231; 2.E.230.236; 2.E.230.237; 2.E.230.238; 2.E.230.239; 2.E.230.154;
2.E.230.157; 2.E.230.166; 2.E.230.169; 2.E.230.172; 2.E.230.175; 2.E.230.240;
2.E.230.244; 2.E.231.228; 2.E.231.229; 2.E.231.230; 2.E.231.231; 2.E.231.236;
2.E.231.237; 2.E.231.238; 2.E.231.239; 2.E.231.154; 2.E.231.157; 2.E.231.166;
2.E.231.169; 2.E.231.172; 2.E.231.175; 2.E.231.240; 2.E.231.244; 2.E.236.228;
10 2.E.236.229; 2.E.236.230; 2.E.236.231; 2.E.236.236; 2.E.236.237; 2.E.236.238;
2.E.236.239; 2.E.236.154; 2.E.236.157; 2.E.236.166; 2.E.236.169; 2.E.236.172;
2.E.236.175; 2.E.236.240; 2.E.236.244; 2.E.237.228; 2.E.237.229; 2.E.237.230;
2.E.237.231; 2.E.237.236; 2.E.237.237; 2.E.237.238; 2.E.237.239; 2.E.237.154;
2.E.237.157; 2.E.237.166; 2.E.237.169; 2.E.237.172; 2.E.237.175; 2.E.237.240;
15 2.E.237.244; 2.E.238.228; 2.E.238.229; 2.E.238.230; 2.E.238.231; 2.E.238.236;
2.E.238.237; 2.E.238.238; 2.E.238.239; 2.E.238.154; 2.E.238.157; 2.E.238.166;
2.E.238.169; 2.E.238.172; 2.E.238.175; 2.E.238.240; 2.E.238.244; 2.E.239.228;
2.E.239.229; 2.E.239.230; 2.E.239.231; 2.E.239.236; 2.E.239.237; 2.E.239.238;
2.E.239.239; 2.E.239.154; 2.E.239.157; 2.E.239.166; 2.E.239.169; 2.E.239.172;
20 2.E.239.175; 2.E.239.240; 2.E.239.244; 2.E.154.228; 2.E.154.229; 2.E.154.230;
2.E.154.231; 2.E.154.236; 2.E.154.237; 2.E.154.238; 2.E.154.239; 2.E.154.154;
2.E.154.157; 2.E.154.166; 2.E.154.169; 2.E.154.172; 2.E.154.175; 2.E.154.240;
2.E.154.244; 2.E.157.228; 2.E.157.229; 2.E.157.230; 2.E.157.231; 2.E.157.236;
2.E.157.237; 2.E.157.238; 2.E.157.239; 2.E.157.154; 2.E.157.157; 2.E.157.166;
25 2.E.157.169; 2.E.157.172; 2.E.157.175; 2.E.157.240; 2.E.157.244; 2.E.166.228;
2.E.166.229; 2.E.166.230; 2.E.166.231; 2.E.166.236; 2.E.166.237; 2.E.166.238;
2.E.166.239; 2.E.166.154; 2.E.166.157; 2.E.166.166; 2.E.166.169; 2.E.166.172;
2.E.166.175; 2.E.166.240; 2.E.166.244; 2.E.169.228; 2.E.169.229; 2.E.169.230;
2.E.169.231; 2.E.169.236; 2.E.169.237; 2.E.169.238; 2.E.169.239; 2.E.169.154;
30 2.E.169.157; 2.E.169.166; 2.E.169.169; 2.E.169.172; 2.E.169.175; 2.E.169.240;

2.E.169.244; 2.E.172.228; 2.E.172.229; 2.E.172.230; 2.E.172.231; 2.E.172.236;
2.E.172.237; 2.E.172.238; 2.E.172.239; 2.E.172.154; 2.E.172.157; 2.E.172.166;
2.E.172.169; 2.E.172.172; 2.E.172.175; 2.E.172.240; 2.E.172.244; 2.E.175.228;
2.E.175.229; 2.E.175.230; 2.E.175.231; 2.E.175.236; 2.E.175.237; 2.E.175.238;
5 2.E.175.239; 2.E.175.154; 2.E.175.157; 2.E.175.166; 2.E.175.169; 2.E.175.172;
2.E.175.175; 2.E.175.240; 2.E.175.244; 2.E.240.228; 2.E.240.229; 2.E.240.230;
2.E.240.231; 2.E.240.236; 2.E.240.237; 2.E.240.238; 2.E.240.239; 2.E.240.154;
2.E.240.157; 2.E.240.166; 2.E.240.169; 2.E.240.172; 2.E.240.175; 2.E.240.240;
2.E.240.244; 2.E.244.228; 2.E.244.229; 2.E.244.230; 2.E.244.231; 2.E.244.236;
10 2.E.244.237; 2.E.244.238; 2.E.244.239; 2.E.244.154; 2.E.244.157; 2.E.244.166;
2.E.244.169; 2.E.244.172; 2.E.244.175; 2.E.244.240; 2.E.244.244;

Prodrugs of 2.G

2.G.228.228; 2.G.228.229; 2.G.228.230; 2.G.228.231; 2.G.228.236;
15 2.G.228.237; 2.G.228.238; 2.G.228.239; 2.G.228.154; 2.G.228.157;
2.G.228.166; 2.G.228.169; 2.G.228.172; 2.G.228.175; 2.G.228.240;
2.G.228.244; 2.G.229.228; 2.G.229.229; 2.G.229.230; 2.G.229.231;
2.G.229.236; 2.G.229.237; 2.G.229.238; 2.G.229.239; 2.G.229.154;
2.G.229.157; 2.G.229.166; 2.G.229.169; 2.G.229.172; 2.G.229.175;
20 2.G.229.240; 2.G.229.244; 2.G.230.228; 2.G.230.229; 2.G.230.230;
2.G.230.231; 2.G.230.236; 2.G.230.237; 2.G.230.238; 2.G.230.239;
2.G.230.154; 2.G.230.157; 2.G.230.166; 2.G.230.169; 2.G.230.172;
2.G.230.175; 2.G.230.240; 2.G.230.244; 2.G.231.228; 2.G.231.229;
2.G.231.230; 2.G.231.231; 2.G.231.236; 2.G.231.237; 2.G.231.238;
25 2.G.231.239; 2.G.231.154; 2.G.231.157; 2.G.231.166; 2.G.231.169;
2.G.231.172; 2.G.231.175; 2.G.231.240; 2.G.231.244; 2.G.236.228;
2.G.236.229; 2.G.236.230; 2.G.236.231; 2.G.236.236; 2.G.236.237;
2.G.236.238; 2.G.236.239; 2.G.236.154; 2.G.236.157; 2.G.236.166;
2.G.236.169; 2.G.236.172; 2.G.236.175; 2.G.236.240; 2.G.236.244;
30 2.G.237.228; 2.G.237.229; 2.G.237.230; 2.G.237.231; 2.G.237.236;

2.G.237.237; 2.G.237.238; 2.G.237.239; 2.G.237.154; 2.G.237.157;
2.G.237.166; 2.G.237.169; 2.G.237.172; 2.G.237.175; 2.G.237.240;
2.G.237.244; 2.G.238.228; 2.G.238.229; 2.G.238.230; 2.G.238.231;
2.G.238.236; 2.G.238.237; 2.G.238.238; 2.G.238.239; 2.G.238.154;
5 2.G.238.157; 2.G.238.166; 2.G.238.169; 2.G.238.172; 2.G.238.175;
2.G.238.240; 2.G.238.244; 2.G.239.228; 2.G.239.229; 2.G.239.230;
2.G.239.231; 2.G.239.236; 2.G.239.237; 2.G.239.238; 2.G.239.239;
2.G.239.154; 2.G.239.157; 2.G.239.166; 2.G.239.169; 2.G.239.172;
2.G.239.175; 2.G.239.240; 2.G.239.244; 2.G.154.228; 2.G.154.229;
10 2.G.154.230; 2.G.154.231; 2.G.154.236; 2.G.154.237; 2.G.154.238;
2.G.154.239; 2.G.154.154; 2.G.154.157; 2.G.154.166; 2.G.154.169;
2.G.154.172; 2.G.154.175; 2.G.154.240; 2.G.154.244; 2.G.157.228;
2.G.157.229; 2.G.157.230; 2.G.157.231; 2.G.157.236; 2.G.157.237;
2.G.157.238; 2.G.157.239; 2.G.157.154; 2.G.157.157; 2.G.157.166;
15 2.G.157.169; 2.G.157.172; 2.G.157.175; 2.G.157.240; 2.G.157.244;
2.G.166.228; 2.G.166.229; 2.G.166.230; 2.G.166.231; 2.G.166.236;
2.G.166.237; 2.G.166.238; 2.G.166.239; 2.G.166.154; 2.G.166.157;
2.G.166.166; 2.G.166.169; 2.G.166.172; 2.G.166.175; 2.G.166.240;
2.G.166.244; 2.G.169.228; 2.G.169.229; 2.G.169.230; 2.G.169.231;
20 2.G.169.236; 2.G.169.237; 2.G.169.238; 2.G.169.239; 2.G.169.154;
2.G.169.157; 2.G.169.166; 2.G.169.169; 2.G.169.172; 2.G.169.175;
2.G.169.240; 2.G.169.244; 2.G.172.228; 2.G.172.229; 2.G.172.230;
2.G.172.231; 2.G.172.236; 2.G.172.237; 2.G.172.238; 2.G.172.239;
2.G.172.154; 2.G.172.157; 2.G.172.166; 2.G.172.169; 2.G.172.172;
25 2.G.172.175; 2.G.172.240; 2.G.172.244; 2.G.175.228; 2.G.175.229;
2.G.175.230; 2.G.175.231; 2.G.175.236; 2.G.175.237; 2.G.175.238;
2.G.175.239; 2.G.175.154; 2.G.175.157; 2.G.175.166; 2.G.175.169;
2.G.175.172; 2.G.175.175; 2.G.175.240; 2.G.175.244; 2.G.240.228;
2.G.240.229; 2.G.240.230; 2.G.240.231; 2.G.240.236; 2.G.240.237;
30 2.G.240.238; 2.G.240.239; 2.G.240.154; 2.G.240.157; 2.G.240.166;

2.G.240.169; 2.G.240.172; 2.G.240.175; 2.G.240.240; 2.G.240.244;
2.G.244.228; 2.G.244.229; 2.G.244.230; 2.G.244.231; 2.G.244.236;
2.G.244.237; 2.G.244.238; 2.G.244.239; 2.G.244.154; 2.G.244.157;
2.G.244.166; 2.G.244.169; 2.G.244.172; 2.G.244.175; 2.G.244.240;
5 2.G.244.244;

Prodrugs of 2.I

2.I.228.228; 2.I.228.229; 2.I.228.230; 2.I.228.231; 2.I.228.236; 2.I.228.237;
2.I.228.238; 2.I.228.239; 2.I.228.154; 2.I.228.157; 2.I.228.166; 2.I.228.169;
10 2.I.228.172; 2.I.228.175; 2.I.228.240; 2.I.228.244; 2.I.229.228; 2.I.229.229;
2.I.229.230; 2.I.229.231; 2.I.229.236; 2.I.229.237; 2.I.229.238; 2.I.229.239;
2.I.229.154; 2.I.229.157; 2.I.229.166; 2.I.229.169; 2.I.229.172; 2.I.229.175;
2.I.229.240; 2.I.229.244; 2.I.230.228; 2.I.230.229; 2.I.230.230; 2.I.230.231;
2.I.230.236; 2.I.230.237; 2.I.230.238; 2.I.230.239; 2.I.230.154; 2.I.230.157;
15 2.I.230.166; 2.I.230.169; 2.I.230.172; 2.I.230.175; 2.I.230.240; 2.I.230.244;
2.I.231.228; 2.I.231.229; 2.I.231.230; 2.I.231.231; 2.I.231.236; 2.I.231.237;
2.I.231.238; 2.I.231.239; 2.I.231.154; 2.I.231.157; 2.I.231.166; 2.I.231.169;
2.I.231.172; 2.I.231.175; 2.I.231.240; 2.I.231.244; 2.I.236.228; 2.I.236.229;
2.I.236.230; 2.I.236.231; 2.I.236.236; 2.I.236.237; 2.I.236.238; 2.I.236.239;
20 2.I.236.154; 2.I.236.157; 2.I.236.166; 2.I.236.169; 2.I.236.172; 2.I.236.175;
2.I.236.240; 2.I.236.244; 2.I.237.228; 2.I.237.229; 2.I.237.230; 2.I.237.231;
2.I.237.236; 2.I.237.237; 2.I.237.238; 2.I.237.239; 2.I.237.154; 2.I.237.157;
2.I.237.166; 2.I.237.169; 2.I.237.172; 2.I.237.175; 2.I.237.240; 2.I.237.244;
2.I.238.228; 2.I.238.229; 2.I.238.230; 2.I.238.231; 2.I.238.236; 2.I.238.237;
25 2.I.238.238; 2.I.238.239; 2.I.238.154; 2.I.238.157; 2.I.238.166; 2.I.238.169;
2.I.238.172; 2.I.238.175; 2.I.238.240; 2.I.238.244; 2.I.239.228; 2.I.239.229;
2.I.239.230; 2.I.239.231; 2.I.239.236; 2.I.239.237; 2.I.239.238; 2.I.239.239;
2.I.239.154; 2.I.239.157; 2.I.239.166; 2.I.239.169; 2.I.239.172; 2.I.239.175;
2.I.239.240; 2.I.239.244; 2.I.154.228; 2.I.154.229; 2.I.154.230; 2.I.154.231;
30 2.I.154.236; 2.I.154.237; 2.I.154.238; 2.I.154.239; 2.I.154.154; 2.I.154.157;

- 2.I.154.166; 2.I.154.169; 2.I.154.172; 2.I.154.175; 2.I.154.240; 2.I.154.244;
2.I.157.228; 2.I.157.229; 2.I.157.230; 2.I.157.231; 2.I.157.236; 2.I.157.237;
2.I.157.238; 2.I.157.239; 2.I.157.154; 2.I.157.157; 2.I.157.166; 2.I.157.169;
2.I.157.172; 2.I.157.175; 2.I.157.240; 2.I.157.244; 2.I.166.228; 2.I.166.229;
5 2.I.166.230; 2.I.166.231; 2.I.166.236; 2.I.166.237; 2.I.166.238; 2.I.166.239;
2.I.166.154; 2.I.166.157; 2.I.166.166; 2.I.166.169; 2.I.166.172; 2.I.166.175;
2.I.166.240; 2.I.166.244; 2.I.169.228; 2.I.169.229; 2.I.169.230; 2.I.169.231;
2.I.169.236; 2.I.169.237; 2.I.169.238; 2.I.169.239; 2.I.169.154; 2.I.169.157;
2.I.169.166; 2.I.169.169; 2.I.169.172; 2.I.169.175; 2.I.169.240; 2.I.169.244;
10 2.I.172.228; 2.I.172.229; 2.I.172.230; 2.I.172.231; 2.I.172.236; 2.I.172.237;
2.I.172.238; 2.I.172.239; 2.I.172.154; 2.I.172.157; 2.I.172.166; 2.I.172.169;
2.I.172.172; 2.I.172.175; 2.I.172.240; 2.I.172.244; 2.I.175.228; 2.I.175.229;
2.I.175.230; 2.I.175.231; 2.I.175.236; 2.I.175.237; 2.I.175.238; 2.I.175.239;
2.I.175.154; 2.I.175.157; 2.I.175.166; 2.I.175.169; 2.I.175.172; 2.I.175.175;
15 2.I.175.240; 2.I.175.244; 2.I.240.228; 2.I.240.229; 2.I.240.230; 2.I.240.231;
2.I.240.236; 2.I.240.237; 2.I.240.238; 2.I.240.239; 2.I.240.154; 2.I.240.157;
2.I.240.166; 2.I.240.169; 2.I.240.172; 2.I.240.175; 2.I.240.240; 2.I.240.244;
2.I.244.228; 2.I.244.229; 2.I.244.230; 2.I.244.231; 2.I.244.236; 2.I.244.237;
2.I.244.238; 2.I.244.239; 2.I.244.154; 2.I.244.157; 2.I.244.166; 2.I.244.169;
20 2.I.244.172; 2.I.244.175; 2.I.244.240; 2.I.244.244;

Prodrugs of 2.J

- 2.J.228.228; 2.J.228.229; 2.J.228.230; 2.J.228.231; 2.J.228.236; 2.J.228.237;
2.J.228.238; 2.J.228.239; 2.J.228.154; 2.J.228.157; 2.J.228.166; 2.J.228.169;
25 2.J.228.172; 2.J.228.175; 2.J.228.240; 2.J.228.244; 2.J.229.228; 2.J.229.229;
2.J.229.230; 2.J.229.231; 2.J.229.236; 2.J.229.237; 2.J.229.238; 2.J.229.239;
2.J.229.154; 2.J.229.157; 2.J.229.166; 2.J.229.169; 2.J.229.172; 2.J.229.175;
2.J.229.240; 2.J.229.244; 2.J.230.228; 2.J.230.229; 2.J.230.230; 2.J.230.231;
2.J.230.236; 2.J.230.237; 2.J.230.238; 2.J.230.239; 2.J.230.154; 2.J.230.157;
30 2.J.230.166; 2.J.230.169; 2.J.230.172; 2.J.230.175; 2.J.230.240; 2.J.230.244;

2.J.231.228; 2.J.231.229; 2.J.231.230; 2.J.231.231; 2.J.231.236; 2.J.231.237;
2.J.231.238; 2.J.231.239; 2.J.231.154; 2.J.231.157; 2.J.231.166; 2.J.231.169;
2.J.231.172; 2.J.231.175; 2.J.231.240; 2.J.231.244; 2.J.236.228; 2.J.236.229;
2.J.236.230; 2.J.236.231; 2.J.236.236; 2.J.236.237; 2.J.236.238; 2.J.236.239;
5 2.J.236.154; 2.J.236.157; 2.J.236.166; 2.J.236.169; 2.J.236.172; 2.J.236.175;
2.J.236.240; 2.J.236.244; 2.J.237.228; 2.J.237.229; 2.J.237.230; 2.J.237.231;
2.J.237.236; 2.J.237.237; 2.J.237.238; 2.J.237.239; 2.J.237.154; 2.J.237.157;
2.J.237.166; 2.J.237.169; 2.J.237.172; 2.J.237.175; 2.J.237.240; 2.J.237.244;
2.J.238.228; 2.J.238.229; 2.J.238.230; 2.J.238.231; 2.J.238.236; 2.J.238.237;
10 2.J.238.238; 2.J.238.239; 2.J.238.154; 2.J.238.157; 2.J.238.166; 2.J.238.169;
2.J.238.172; 2.J.238.175; 2.J.238.240; 2.J.238.244; 2.J.239.228; 2.J.239.229;
2.J.239.230; 2.J.239.231; 2.J.239.236; 2.J.239.237; 2.J.239.238; 2.J.239.239;
2.J.239.154; 2.J.239.157; 2.J.239.166; 2.J.239.169; 2.J.239.172; 2.J.239.175;
2.J.239.240; 2.J.239.244; 2.J.154.228; 2.J.154.229; 2.J.154.230; 2.J.154.231;
15 2.J.154.236; 2.J.154.237; 2.J.154.238; 2.J.154.239; 2.J.154.154; 2.J.154.157;
2.J.154.166; 2.J.154.169; 2.J.154.172; 2.J.154.175; 2.J.154.240; 2.J.154.244;
2.J.157.228; 2.J.157.229; 2.J.157.230; 2.J.157.231; 2.J.157.236; 2.J.157.237;
2.J.157.238; 2.J.157.239; 2.J.157.154; 2.J.157.157; 2.J.157.166; 2.J.157.169;
2.J.157.172; 2.J.157.175; 2.J.157.240; 2.J.157.244; 2.J.166.228; 2.J.166.229;
20 2.J.166.230; 2.J.166.231; 2.J.166.236; 2.J.166.237; 2.J.166.238; 2.J.166.239;
2.J.166.154; 2.J.166.157; 2.J.166.166; 2.J.166.169; 2.J.166.172; 2.J.166.175;
2.J.166.240; 2.J.166.244; 2.J.169.228; 2.J.169.229; 2.J.169.230; 2.J.169.231;
2.J.169.236; 2.J.169.237; 2.J.169.238; 2.J.169.239; 2.J.169.154; 2.J.169.157;
2.J.169.166; 2.J.169.169; 2.J.169.172; 2.J.169.175; 2.J.169.240; 2.J.169.244;
25 2.J.172.228; 2.J.172.229; 2.J.172.230; 2.J.172.231; 2.J.172.236; 2.J.172.237;
2.J.172.238; 2.J.172.239; 2.J.172.154; 2.J.172.157; 2.J.172.166; 2.J.172.169;
2.J.172.172; 2.J.172.175; 2.J.172.240; 2.J.172.244; 2.J.175.228; 2.J.175.229;
2.J.175.230; 2.J.175.231; 2.J.175.236; 2.J.175.237; 2.J.175.238; 2.J.175.239;
2.J.175.154; 2.J.175.157; 2.J.175.166; 2.J.175.169; 2.J.175.172; 2.J.175.175;
30 2.J.175.240; 2.J.175.244; 2.J.240.228; 2.J.240.229; 2.J.240.230; 2.J.240.231;

2.J.240.236; 2.J.240.237; 2.J.240.238; 2.J.240.239; 2.J.240.154; 2.J.240.157;
2.J.240.166; 2.J.240.169; 2.J.240.172; 2.J.240.175; 2.J.240.240; 2.J.240.244;
2.J.244.228; 2.J.244.229; 2.J.244.230; 2.J.244.231; 2.J.244.236; 2.J.244.237;
2.J.244.238; 2.J.244.239; 2.J.244.154; 2.J.244.157; 2.J.244.166; 2.J.244.169;
5 2.J.244.172; 2.J.244.175; 2.J.244.240; 2.J.244.244;

Prodrugs of 2.L

2.L.228.228; 2.L.228.229; 2.L.228.230; 2.L.228.231; 2.L.228.236;
2.L.228.237; 2.L.228.238; 2.L.228.239; 2.L.228.154; 2.L.228.157; 2.L.228.166;
10 2.L.228.169; 2.L.228.172; 2.L.228.175; 2.L.228.240; 2.L.228.244; 2.L.229.228;
2.L.229.229; 2.L.229.230; 2.L.229.231; 2.L.229.236; 2.L.229.237; 2.L.229.238;
2.L.229.239; 2.L.229.154; 2.L.229.157; 2.L.229.166; 2.L.229.169; 2.L.229.172;
2.L.229.175; 2.L.229.240; 2.L.229.244; 2.L.230.228; 2.L.230.229; 2.L.230.230;
2.L.230.231; 2.L.230.236; 2.L.230.237; 2.L.230.238; 2.L.230.239; 2.L.230.154;
15 2.L.230.157; 2.L.230.166; 2.L.230.169; 2.L.230.172; 2.L.230.175; 2.L.230.240;
2.L.230.244; 2.L.231.228; 2.L.231.229; 2.L.231.230; 2.L.231.231; 2.L.231.236;
2.L.231.237; 2.L.231.238; 2.L.231.239; 2.L.231.154; 2.L.231.157; 2.L.231.166;
2.L.231.169; 2.L.231.172; 2.L.231.175; 2.L.231.240; 2.L.231.244; 2.L.236.228;
2.L.236.229; 2.L.236.230; 2.L.236.231; 2.L.236.236; 2.L.236.237; 2.L.236.238;
20 2.L.236.239; 2.L.236.154; 2.L.236.157; 2.L.236.166; 2.L.236.169; 2.L.236.172;
2.L.236.175; 2.L.236.240; 2.L.236.244; 2.L.237.228; 2.L.237.229; 2.L.237.230;
2.L.237.231; 2.L.237.236; 2.L.237.237; 2.L.237.238; 2.L.237.239; 2.L.237.154;
2.L.237.157; 2.L.237.166; 2.L.237.169; 2.L.237.172; 2.L.237.175; 2.L.237.240;
2.L.237.244; 2.L.238.228; 2.L.238.229; 2.L.238.230; 2.L.238.231; 2.L.238.236;
25 2.L.238.237; 2.L.238.238; 2.L.238.239; 2.L.238.154; 2.L.238.157; 2.L.238.166;
2.L.238.169; 2.L.238.172; 2.L.238.175; 2.L.238.240; 2.L.238.244; 2.L.239.228;
2.L.239.229; 2.L.239.230; 2.L.239.231; 2.L.239.236; 2.L.239.237; 2.L.239.238;
2.L.239.239; 2.L.239.154; 2.L.239.157; 2.L.239.166; 2.L.239.169; 2.L.239.172;
2.L.239.175; 2.L.239.240; 2.L.239.244; 2.L.154.228; 2.L.154.229; 2.L.154.230;
30 2.L.154.231; 2.L.154.236; 2.L.154.237; 2.L.154.238; 2.L.154.239; 2.L.154.154;

2.L.154.157; 2.L.154.166; 2.L.154.169; 2.L.154.172; 2.L.154.175; 2.L.154.240;
 2.L.154.244; 2.L.157.228; 2.L.157.229; 2.L.157.230; 2.L.157.231; 2.L.157.236;
 2.L.157.237; 2.L.157.238; 2.L.157.239; 2.L.157.154; 2.L.157.157; 2.L.157.166;
 2.L.157.169; 2.L.157.172; 2.L.157.175; 2.L.157.240; 2.L.157.244; 2.L.166.228;
 5 2.L.166.229; 2.L.166.230; 2.L.166.231; 2.L.166.236; 2.L.166.237; 2.L.166.238;
 2.L.166.239; 2.L.166.154; 2.L.166.157; 2.L.166.166; 2.L.166.169; 2.L.166.172;
 2.L.166.175; 2.L.166.240; 2.L.166.244; 2.L.169.228; 2.L.169.229; 2.L.169.230;
 2.L.169.231; 2.L.169.236; 2.L.169.237; 2.L.169.238; 2.L.169.239; 2.L.169.154;
 2.L.169.157; 2.L.169.166; 2.L.169.169; 2.L.169.172; 2.L.169.175; 2.L.169.240;
 10 2.L.169.244; 2.L.172.228; 2.L.172.229; 2.L.172.230; 2.L.172.231; 2.L.172.236;
 2.L.172.237; 2.L.172.238; 2.L.172.239; 2.L.172.154; 2.L.172.157; 2.L.172.166;
 2.L.172.169; 2.L.172.172; 2.L.172.175; 2.L.172.240; 2.L.172.244; 2.L.175.228;
 2.L.175.229; 2.L.175.230; 2.L.175.231; 2.L.175.236; 2.L.175.237; 2.L.175.238;
 2.L.175.239; 2.L.175.154; 2.L.175.157; 2.L.175.166; 2.L.175.169; 2.L.175.172;
 15 2.L.175.175; 2.L.175.240; 2.L.175.244; 2.L.240.228; 2.L.240.229; 2.L.240.230;
 2.L.240.231; 2.L.240.236; 2.L.240.237; 2.L.240.238; 2.L.240.239; 2.L.240.154;
 2.L.240.157; 2.L.240.166; 2.L.240.169; 2.L.240.172; 2.L.240.175; 2.L.240.240;
 2.L.240.244; 2.L.244.228; 2.L.244.229; 2.L.244.230; 2.L.244.231; 2.L.244.236;
 2.L.244.237; 2.L.244.238; 2.L.244.239; 2.L.244.154; 2.L.244.157; 2.L.244.166;
 20 2.L.244.169; 2.L.244.172; 2.L.244.175; 2.L.244.240; 2.L.244.244;

Prodrugs of 2.O

2.O.228.228; 2.O.228.229; 2.O.228.230; 2.O.228.231; 2.O.228.236;
 2.O.228.237; 2.O.228.238; 2.O.228.239; 2.O.228.154; 2.O.228.157;
 25 2.O.228.166; 2.O.228.169; 2.O.228.172; 2.O.228.175; 2.O.228.240;
 2.O.228.244; 2.O.229.228; 2.O.229.229; 2.O.229.230; 2.O.229.231;
 2.O.229.236; 2.O.229.237; 2.O.229.238; 2.O.229.239; 2.O.229.154;
 2.O.229.157; 2.O.229.166; 2.O.229.169; 2.O.229.172; 2.O.229.175;
 2.O.229.240; 2.O.229.244; 2.O.230.228; 2.O.230.229; 2.O.230.230;
 30 2.O.230.231; 2.O.230.236; 2.O.230.237; 2.O.230.238; 2.O.230.239;

2.O.230.154; 2.O.230.157; 2.O.230.166; 2.O.230.169; 2.O.230.172;
2.O.230.175; 2.O.230.240; 2.O.230.244; 2.O.231.228; 2.O.231.229;
2.O.231.230; 2.O.231.231; 2.O.231.236; 2.O.231.237; 2.O.231.238;
2.O.231.239; 2.O.231.154; 2.O.231.157; 2.O.231.166; 2.O.231.169;
5 2.O.231.172; 2.O.231.175; 2.O.231.240; 2.O.231.244; 2.O.236.228;
2.O.236.229; 2.O.236.230; 2.O.236.231; 2.O.236.236; 2.O.236.237;
2.O.236.238; 2.O.236.239; 2.O.236.154; 2.O.236.157; 2.O.236.166;
2.O.236.169; 2.O.236.172; 2.O.236.175; 2.O.236.240; 2.O.236.244;
2.O.237.228; 2.O.237.229; 2.O.237.230; 2.O.237.231; 2.O.237.236;
10 2.O.237.237; 2.O.237.238; 2.O.237.239; 2.O.237.154; 2.O.237.157;
2.O.237.166; 2.O.237.169; 2.O.237.172; 2.O.237.175; 2.O.237.240;
2.O.237.244; 2.O.238.228; 2.O.238.229; 2.O.238.230; 2.O.238.231;
2.O.238.236; 2.O.238.237; 2.O.238.238; 2.O.238.239; 2.O.238.154;
2.O.238.157; 2.O.238.166; 2.O.238.169; 2.O.238.172; 2.O.238.175;
15 2.O.238.240; 2.O.238.244; 2.O.239.228; 2.O.239.229; 2.O.239.230;
2.O.239.231; 2.O.239.236; 2.O.239.237; 2.O.239.238; 2.O.239.239;
2.O.239.154; 2.O.239.157; 2.O.239.166; 2.O.239.169; 2.O.239.172;
2.O.239.175; 2.O.239.240; 2.O.239.244; 2.O.154.228; 2.O.154.229;
2.O.154.230; 2.O.154.231; 2.O.154.236; 2.O.154.237; 2.O.154.238;
20 2.O.154.239; 2.O.154.154; 2.O.154.157; 2.O.154.166; 2.O.154.169;
2.O.154.172; 2.O.154.175; 2.O.154.240; 2.O.154.244; 2.O.157.228;
2.O.157.229; 2.O.157.230; 2.O.157.231; 2.O.157.236; 2.O.157.237;
2.O.157.238; 2.O.157.239; 2.O.157.154; 2.O.157.157; 2.O.157.166;
2.O.157.169; 2.O.157.172; 2.O.157.175; 2.O.157.240; 2.O.157.244;
25 2.O.166.228; 2.O.166.229; 2.O.166.230; 2.O.166.231; 2.O.166.236;
2.O.166.237; 2.O.166.238; 2.O.166.239; 2.O.166.154; 2.O.166.157;
2.O.166.166; 2.O.166.169; 2.O.166.172; 2.O.166.175; 2.O.166.240;
2.O.166.244; 2.O.169.228; 2.O.169.229; 2.O.169.230; 2.O.169.231;
2.O.169.236; 2.O.169.237; 2.O.169.238; 2.O.169.239; 2.O.169.154;
30 2.O.169.157; 2.O.169.166; 2.O.169.169; 2.O.169.172; 2.O.169.175;

2.O.169.240; 2.O.169.244; 2.O.172.228; 2.O.172.229; 2.O.172.230;
2.O.172.231; 2.O.172.236; 2.O.172.237; 2.O.172.238; 2.O.172.239;
2.O.172.154; 2.O.172.157; 2.O.172.166; 2.O.172.169; 2.O.172.172;
2.O.172.175; 2.O.172.240; 2.O.172.244; 2.O.175.228; 2.O.175.229;
5 2.O.175.230; 2.O.175.231; 2.O.175.236; 2.O.175.237; 2.O.175.238;
2.O.175.239; 2.O.175.154; 2.O.175.157; 2.O.175.166; 2.O.175.169;
2.O.175.172; 2.O.175.175; 2.O.175.240; 2.O.175.244; 2.O.240.228;
2.O.240.229; 2.O.240.230; 2.O.240.231; 2.O.240.236; 2.O.240.237;
2.O.240.238; 2.O.240.239; 2.O.240.154; 2.O.240.157; 2.O.240.166;
10 2.O.240.169; 2.O.240.172; 2.O.240.175; 2.O.240.240; 2.O.240.244;
2.O.244.228; 2.O.244.229; 2.O.244.230; 2.O.244.231; 2.O.244.236;
2.O.244.237; 2.O.244.238; 2.O.244.239; 2.O.244.154; 2.O.244.157;
2.O.244.166; 2.O.244.169; 2.O.244.172; 2.O.244.175; 2.O.244.240;
2.O.244.244;

15

Prodrugs of 2.P

2.P.228.228; 2.P.228.229; 2.P.228.230; 2.P.228.231; 2.P.228.236;
2.P.228.237; 2.P.228.238; 2.P.228.239; 2.P.228.154; 2.P.228.157; 2.P.228.166;
2.P.228.169; 2.P.228.172; 2.P.228.175; 2.P.228.240; 2.P.228.244; 2.P.229.228;
20 2.P.229.229; 2.P.229.230; 2.P.229.231; 2.P.229.236; 2.P.229.237; 2.P.229.238;
2.P.229.239; 2.P.229.154; 2.P.229.157; 2.P.229.166; 2.P.229.169; 2.P.229.172;
2.P.229.175; 2.P.229.240; 2.P.229.244; 2.P.230.228; 2.P.230.229; 2.P.230.230;
2.P.230.231; 2.P.230.236; 2.P.230.237; 2.P.230.238; 2.P.230.239; 2.P.230.154;
2.P.230.157; 2.P.230.166; 2.P.230.169; 2.P.230.172; 2.P.230.175; 2.P.230.240;
25 2.P.230.244; 2.P.231.228; 2.P.231.229; 2.P.231.230; 2.P.231.231; 2.P.231.236;
2.P.231.237; 2.P.231.238; 2.P.231.239; 2.P.231.154; 2.P.231.157; 2.P.231.166;
2.P.231.169; 2.P.231.172; 2.P.231.175; 2.P.231.240; 2.P.231.244; 2.P.236.228;
2.P.236.229; 2.P.236.230; 2.P.236.231; 2.P.236.236; 2.P.236.237; 2.P.236.238;
2.P.236.239; 2.P.236.154; 2.P.236.157; 2.P.236.166; 2.P.236.169; 2.P.236.172;
30 2.P.236.175; 2.P.236.240; 2.P.236.244; 2.P.237.228; 2.P.237.229; 2.P.237.230;

2.P.237.231; 2.P.237.236; 2.P.237.237; 2.P.237.238; 2.P.237.239; 2.P.237.154;
2.P.237.157; 2.P.237.166; 2.P.237.169; 2.P.237.172; 2.P.237.175; 2.P.237.240;
2.P.237.244; 2.P.238.228; 2.P.238.229; 2.P.238.230; 2.P.238.231; 2.P.238.236;
2.P.238.237; 2.P.238.238; 2.P.238.239; 2.P.238.154; 2.P.238.157; 2.P.238.166;
5 2.P.238.169; 2.P.238.172; 2.P.238.175; 2.P.238.240; 2.P.238.244; 2.P.239.228;
2.P.239.229; 2.P.239.230; 2.P.239.231; 2.P.239.236; 2.P.239.237; 2.P.239.238;
2.P.239.239; 2.P.239.154; 2.P.239.157; 2.P.239.166; 2.P.239.169; 2.P.239.172;
2.P.239.175; 2.P.239.240; 2.P.239.244; 2.P.154.228; 2.P.154.229; 2.P.154.230;
2.P.154.231; 2.P.154.236; 2.P.154.237; 2.P.154.238; 2.P.154.239; 2.P.154.154;
10 2.P.154.157; 2.P.154.166; 2.P.154.169; 2.P.154.172; 2.P.154.175; 2.P.154.240;
2.P.154.244; 2.P.157.228; 2.P.157.229; 2.P.157.230; 2.P.157.231; 2.P.157.236;
2.P.157.237; 2.P.157.238; 2.P.157.239; 2.P.157.154; 2.P.157.157; 2.P.157.166;
2.P.157.169; 2.P.157.172; 2.P.157.175; 2.P.157.240; 2.P.157.244; 2.P.166.228;
2.P.166.229; 2.P.166.230; 2.P.166.231; 2.P.166.236; 2.P.166.237; 2.P.166.238;
15 2.P.166.239; 2.P.166.154; 2.P.166.157; 2.P.166.166; 2.P.166.169; 2.P.166.172;
2.P.166.175; 2.P.166.240; 2.P.166.244; 2.P.169.228; 2.P.169.229; 2.P.169.230;
2.P.169.231; 2.P.169.236; 2.P.169.237; 2.P.169.238; 2.P.169.239; 2.P.169.154;
2.P.169.157; 2.P.169.166; 2.P.169.169; 2.P.169.172; 2.P.169.175; 2.P.169.240;
2.P.169.244; 2.P.172.228; 2.P.172.229; 2.P.172.230; 2.P.172.231; 2.P.172.236;
20 2.P.172.237; 2.P.172.238; 2.P.172.239; 2.P.172.154; 2.P.172.157; 2.P.172.166;
2.P.172.169; 2.P.172.172; 2.P.172.175; 2.P.172.240; 2.P.172.244; 2.P.175.228;
2.P.175.229; 2.P.175.230; 2.P.175.231; 2.P.175.236; 2.P.175.237; 2.P.175.238;
2.P.175.239; 2.P.175.154; 2.P.175.157; 2.P.175.166; 2.P.175.169; 2.P.175.172;
2.P.175.175; 2.P.175.240; 2.P.175.244; 2.P.240.228; 2.P.240.229; 2.P.240.230;
25 2.P.240.231; 2.P.240.236; 2.P.240.237; 2.P.240.238; 2.P.240.239; 2.P.240.154;
2.P.240.157; 2.P.240.166; 2.P.240.169; 2.P.240.172; 2.P.240.175; 2.P.240.240;
2.P.240.244; 2.P.244.228; 2.P.244.229; 2.P.244.230; 2.P.244.231; 2.P.244.236;
2.P.244.237; 2.P.244.238; 2.P.244.239; 2.P.244.154; 2.P.244.157; 2.P.244.166;
2.P.244.169; 2.P.244.172; 2.P.244.175; 2.P.244.240; 2.P.244.244;

Prodrugs of 2.U

- 2.U.228.228; 2.U.228.229; 2.U.228.230; 2.U.228.231; 2.U.228.236;
2.U.228.237; 2.U.228.238; 2.U.228.239; 2.U.228.154; 2.U.228.157;
2.U.228.166; 2.U.228.169; 2.U.228.172; 2.U.228.175; 2.U.228.240;
5 2.U.228.244; 2.U.229.228; 2.U.229.229; 2.U.229.230; 2.U.229.231;
2.U.229.236; 2.U.229.237; 2.U.229.238; 2.U.229.239; 2.U.229.154;
2.U.229.157; 2.U.229.166; 2.U.229.169; 2.U.229.172; 2.U.229.175;
2.U.229.240; 2.U.229.244; 2.U.230.228; 2.U.230.229; 2.U.230.230;
2.U.230.231; 2.U.230.236; 2.U.230.237; 2.U.230.238; 2.U.230.239;
10 2.U.230.154; 2.U.230.157; 2.U.230.166; 2.U.230.169; 2.U.230.172;
2.U.230.175; 2.U.230.240; 2.U.230.244; 2.U.231.228; 2.U.231.229;
2.U.231.230; 2.U.231.231; 2.U.231.236; 2.U.231.237; 2.U.231.238;
2.U.231.239; 2.U.231.154; 2.U.231.157; 2.U.231.166; 2.U.231.169;
2.U.231.172; 2.U.231.175; 2.U.231.240; 2.U.231.244; 2.U.236.228;
15 2.U.236.229; 2.U.236.230; 2.U.236.231; 2.U.236.236; 2.U.236.237;
2.U.236.238; 2.U.236.239; 2.U.236.154; 2.U.236.157; 2.U.236.166;
2.U.236.169; 2.U.236.172; 2.U.236.175; 2.U.236.240; 2.U.236.244;
2.U.237.228; 2.U.237.229; 2.U.237.230; 2.U.237.231; 2.U.237.236;
2.U.237.237; 2.U.237.238; 2.U.237.239; 2.U.237.154; 2.U.237.157;
20 2.U.237.166; 2.U.237.169; 2.U.237.172; 2.U.237.175; 2.U.237.240;
2.U.237.244; 2.U.238.228; 2.U.238.229; 2.U.238.230; 2.U.238.231;
2.U.238.236; 2.U.238.237; 2.U.238.238; 2.U.238.239; 2.U.238.154;
2.U.238.157; 2.U.238.166; 2.U.238.169; 2.U.238.172; 2.U.238.175;
2.U.238.240; 2.U.238.244; 2.U.239.228; 2.U.239.229; 2.U.239.230;
25 2.U.239.231; 2.U.239.236; 2.U.239.237; 2.U.239.238; 2.U.239.239;
2.U.239.154; 2.U.239.157; 2.U.239.166; 2.U.239.169; 2.U.239.172;
2.U.239.175; 2.U.239.240; 2.U.239.244; 2.U.154.228; 2.U.154.229;
2.U.154.230; 2.U.154.231; 2.U.154.236; 2.U.154.237; 2.U.154.238;
2.U.154.239; 2.U.154.154; 2.U.154.157; 2.U.154.166; 2.U.154.169;
30 2.U.154.172; 2.U.154.175; 2.U.154.240; 2.U.154.244; 2.U.157.228;

- 2.U.157.229; 2.U.157.230; 2.U.157.231; 2.U.157.236; 2.U.157.237;
2.U.157.238; 2.U.157.239; 2.U.157.154; 2.U.157.157; 2.U.157.166;
2.U.157.169; 2.U.157.172; 2.U.157.175; 2.U.157.240; 2.U.157.244;
2.U.166.228; 2.U.166.229; 2.U.166.230; 2.U.166.231; 2.U.166.236;
5 2.U.166.237; 2.U.166.238; 2.U.166.239; 2.U.166.154; 2.U.166.157;
2.U.166.166; 2.U.166.169; 2.U.166.172; 2.U.166.175; 2.U.166.240;
2.U.166.244; 2.U.169.228; 2.U.169.229; 2.U.169.230; 2.U.169.231;
2.U.169.236; 2.U.169.237; 2.U.169.238; 2.U.169.239; 2.U.169.154;
2.U.169.157; 2.U.169.166; 2.U.169.169; 2.U.169.172; 2.U.169.175;
10 2.U.169.240; 2.U.169.244; 2.U.172.228; 2.U.172.229; 2.U.172.230;
2.U.172.231; 2.U.172.236; 2.U.172.237; 2.U.172.238; 2.U.172.239;
2.U.172.154; 2.U.172.157; 2.U.172.166; 2.U.172.169; 2.U.172.172;
2.U.172.175; 2.U.172.240; 2.U.172.244; 2.U.175.228; 2.U.175.229;
2.U.175.230; 2.U.175.231; 2.U.175.236; 2.U.175.237; 2.U.175.238;
15 2.U.175.239; 2.U.175.154; 2.U.175.157; 2.U.175.166; 2.U.175.169;
2.U.175.172; 2.U.175.175; 2.U.175.240; 2.U.175.244; 2.U.240.228;
2.U.240.229; 2.U.240.230; 2.U.240.231; 2.U.240.236; 2.U.240.237;
2.U.240.238; 2.U.240.239; 2.U.240.154; 2.U.240.157; 2.U.240.166;
2.U.240.169; 2.U.240.172; 2.U.240.175; 2.U.240.240; 2.U.240.244;
20 2.U.244.228; 2.U.244.229; 2.U.244.230; 2.U.244.231; 2.U.244.236;
2.U.244.237; 2.U.244.238; 2.U.244.239; 2.U.244.154; 2.U.244.157;
2.U.244.166; 2.U.244.169; 2.U.244.172; 2.U.244.175; 2.U.244.240;
2.U.244.244;
- 25 Prodrugs of 2.W
2.W.228.228; 2.W.228.229; 2.W.228.230; 2.W.228.231; 2.W.228.236;
2.W.228.237; 2.W.228.238; 2.W.228.239; 2.W.228.154; 2.W.228.157;
2.W.228.166; 2.W.228.169; 2.W.228.172; 2.W.228.175; 2.W.228.240;
2.W.228.244; 2.W.229.228; 2.W.229.229; 2.W.229.230; 2.W.229.231;
30 2.W.229.236; 2.W.229.237; 2.W.229.238; 2.W.229.239; 2.W.229.154;

2.W.229.157; 2.W.229.166; 2.W.229.169; 2.W.229.172; 2.W.229.175;
2.W.229.240; 2.W.229.244; 2.W.230.228; 2.W.230.229; 2.W.230.230;
2.W.230.231; 2.W.230.236; 2.W.230.237; 2.W.230.238; 2.W.230.239;
2.W.230.154; 2.W.230.157; 2.W.230.166; 2.W.230.169; 2.W.230.172;
5 2.W.230.175; 2.W.230.240; 2.W.230.244; 2.W.231.228; 2.W.231.229;
2.W.231.230; 2.W.231.231; 2.W.231.236; 2.W.231.237; 2.W.231.238;
2.W.231.239; 2.W.231.154; 2.W.231.157; 2.W.231.166; 2.W.231.169;
2.W.231.172; 2.W.231.175; 2.W.231.240; 2.W.231.244; 2.W.236.228;
2.W.236.229; 2.W.236.230; 2.W.236.231; 2.W.236.236; 2.W.236.237;
10 2.W.236.238; 2.W.236.239; 2.W.236.154; 2.W.236.157; 2.W.236.166;
2.W.236.169; 2.W.236.172; 2.W.236.175; 2.W.236.240; 2.W.236.244;
2.W.237.228; 2.W.237.229; 2.W.237.230; 2.W.237.231; 2.W.237.236;
2.W.237.237; 2.W.237.238; 2.W.237.239; 2.W.237.154; 2.W.237.157;
2.W.237.166; 2.W.237.169; 2.W.237.172; 2.W.237.175; 2.W.237.240;
15 2.W.237.244; 2.W.238.228; 2.W.238.229; 2.W.238.230; 2.W.238.231;
2.W.238.236; 2.W.238.237; 2.W.238.238; 2.W.238.239; 2.W.238.154;
2.W.238.157; 2.W.238.166; 2.W.238.169; 2.W.238.172; 2.W.238.175;
2.W.238.240; 2.W.238.244; 2.W.239.228; 2.W.239.229; 2.W.239.230;
2.W.239.231; 2.W.239.236; 2.W.239.237; 2.W.239.238; 2.W.239.239;
20 2.W.239.154; 2.W.239.157; 2.W.239.166; 2.W.239.169; 2.W.239.172;
2.W.239.175; 2.W.239.240; 2.W.239.244; 2.W.154.228; 2.W.154.229;
2.W.154.230; 2.W.154.231; 2.W.154.236; 2.W.154.237; 2.W.154.238;
2.W.154.239; 2.W.154.154; 2.W.154.157; 2.W.154.166; 2.W.154.169;
2.W.154.172; 2.W.154.175; 2.W.154.240; 2.W.154.244; 2.W.157.228;
25 2.W.157.229; 2.W.157.230; 2.W.157.231; 2.W.157.236; 2.W.157.237;
2.W.157.238; 2.W.157.239; 2.W.157.154; 2.W.157.157; 2.W.157.166;
2.W.157.169; 2.W.157.172; 2.W.157.175; 2.W.157.240; 2.W.157.244;
2.W.166.228; 2.W.166.229; 2.W.166.230; 2.W.166.231; 2.W.166.236;
2.W.166.237; 2.W.166.238; 2.W.166.239; 2.W.166.154; 2.W.166.157;
30 2.W.166.166; 2.W.166.169; 2.W.166.172; 2.W.166.175; 2.W.166.240;

2.W.166.244; 2.W.169.228; 2.W.169.229; 2.W.169.230; 2.W.169.231;
 2.W.169.236; 2.W.169.237; 2.W.169.238; 2.W.169.239; 2.W.169.154;
 2.W.169.157; 2.W.169.166; 2.W.169.169; 2.W.169.172; 2.W.169.175;
 2.W.169.240; 2.W.169.244; 2.W.172.228; 2.W.172.229; 2.W.172.230;
 5 2.W.172.231; 2.W.172.236; 2.W.172.237; 2.W.172.238; 2.W.172.239;
 2.W.172.154; 2.W.172.157; 2.W.172.166; 2.W.172.169; 2.W.172.172;
 2.W.172.175; 2.W.172.240; 2.W.172.244; 2.W.175.228; 2.W.175.229;
 2.W.175.230; 2.W.175.231; 2.W.175.236; 2.W.175.237; 2.W.175.238;
 2.W.175.239; 2.W.175.154; 2.W.175.157; 2.W.175.166; 2.W.175.169;
 10 2.W.175.172; 2.W.175.175; 2.W.175.240; 2.W.175.244; 2.W.240.228;
 2.W.240.229; 2.W.240.230; 2.W.240.231; 2.W.240.236; 2.W.240.237;
 2.W.240.238; 2.W.240.239; 2.W.240.154; 2.W.240.157; 2.W.240.166;
 2.W.240.169; 2.W.240.172; 2.W.240.175; 2.W.240.240; 2.W.240.244;
 2.W.244.228; 2.W.244.229; 2.W.244.230; 2.W.244.231; 2.W.244.236;
 15 2.W.244.237; 2.W.244.238; 2.W.244.239; 2.W.244.154; 2.W.244.157;
 2.W.244.166; 2.W.244.169; 2.W.244.172; 2.W.244.175; 2.W.244.240;
 2.W.244.244;

Prodrugs of 2.Y

20 2.Y.228.228; 2.Y.228.229; 2.Y.228.230; 2.Y.228.231; 2.Y.228.236;
 2.Y.228.237; 2.Y.228.238; 2.Y.228.239; 2.Y.228.154; 2.Y.228.157; 2.Y.228.166;
 2.Y.228.169; 2.Y.228.172; 2.Y.228.175; 2.Y.228.240; 2.Y.228.244; 2.Y.229.228;
 2.Y.229.229; 2.Y.229.230; 2.Y.229.231; 2.Y.229.236; 2.Y.229.237; 2.Y.229.238;
 2.Y.229.239; 2.Y.229.154; 2.Y.229.157; 2.Y.229.166; 2.Y.229.169; 2.Y.229.172;
 25 2.Y.229.175; 2.Y.229.240; 2.Y.229.244; 2.Y.230.228; 2.Y.230.229; 2.Y.230.230;
 2.Y.230.231; 2.Y.230.236; 2.Y.230.237; 2.Y.230.238; 2.Y.230.239; 2.Y.230.154;
 2.Y.230.157; 2.Y.230.166; 2.Y.230.169; 2.Y.230.172; 2.Y.230.175; 2.Y.230.240;
 2.Y.230.244; 2.Y.231.228; 2.Y.231.229; 2.Y.231.230; 2.Y.231.231; 2.Y.231.236;
 2.Y.231.237; 2.Y.231.238; 2.Y.231.239; 2.Y.231.154; 2.Y.231.157; 2.Y.231.166;
 30 2.Y.231.169; 2.Y.231.172; 2.Y.231.175; 2.Y.231.240; 2.Y.231.244; 2.Y.236.228;

2.Y.236.229; 2.Y.236.230; 2.Y.236.231; 2.Y.236.236; 2.Y.236.237; 2.Y.236.238;
2.Y.236.239; 2.Y.236.154; 2.Y.236.157; 2.Y.236.166; 2.Y.236.169; 2.Y.236.172;
2.Y.236.175; 2.Y.236.240; 2.Y.236.244; 2.Y.237.228; 2.Y.237.229; 2.Y.237.230;
2.Y.237.231; 2.Y.237.236; 2.Y.237.237; 2.Y.237.238; 2.Y.237.239; 2.Y.237.154;
5 2.Y.237.157; 2.Y.237.166; 2.Y.237.169; 2.Y.237.172; 2.Y.237.175; 2.Y.237.240;
2.Y.237.244; 2.Y.238.228; 2.Y.238.229; 2.Y.238.230; 2.Y.238.231; 2.Y.238.236;
2.Y.238.237; 2.Y.238.238; 2.Y.238.239; 2.Y.238.154; 2.Y.238.157; 2.Y.238.166;
2.Y.238.169; 2.Y.238.172; 2.Y.238.175; 2.Y.238.240; 2.Y.238.244; 2.Y.239.228;
2.Y.239.229; 2.Y.239.230; 2.Y.239.231; 2.Y.239.236; 2.Y.239.237; 2.Y.239.238;
10 2.Y.239.239; 2.Y.239.154; 2.Y.239.157; 2.Y.239.166; 2.Y.239.169; 2.Y.239.172;
2.Y.239.175; 2.Y.239.240; 2.Y.239.244; 2.Y.154.228; 2.Y.154.229; 2.Y.154.230;
2.Y.154.231; 2.Y.154.236; 2.Y.154.237; 2.Y.154.238; 2.Y.154.239; 2.Y.154.154;
2.Y.154.157; 2.Y.154.166; 2.Y.154.169; 2.Y.154.172; 2.Y.154.175; 2.Y.154.240;
2.Y.154.244; 2.Y.157.228; 2.Y.157.229; 2.Y.157.230; 2.Y.157.231; 2.Y.157.236;
15 2.Y.157.237; 2.Y.157.238; 2.Y.157.239; 2.Y.157.154; 2.Y.157.157; 2.Y.157.166;
2.Y.157.169; 2.Y.157.172; 2.Y.157.175; 2.Y.157.240; 2.Y.157.244; 2.Y.166.228;
2.Y.166.229; 2.Y.166.230; 2.Y.166.231; 2.Y.166.236; 2.Y.166.237; 2.Y.166.238;
2.Y.166.239; 2.Y.166.154; 2.Y.166.157; 2.Y.166.166; 2.Y.166.169; 2.Y.166.172;
2.Y.166.175; 2.Y.166.240; 2.Y.166.244; 2.Y.169.228; 2.Y.169.229; 2.Y.169.230;
20 2.Y.169.231; 2.Y.169.236; 2.Y.169.237; 2.Y.169.238; 2.Y.169.239; 2.Y.169.154;
2.Y.169.157; 2.Y.169.166; 2.Y.169.169; 2.Y.169.172; 2.Y.169.175; 2.Y.169.240;
2.Y.169.244; 2.Y.172.228; 2.Y.172.229; 2.Y.172.230; 2.Y.172.231; 2.Y.172.236;
2.Y.172.237; 2.Y.172.238; 2.Y.172.239; 2.Y.172.154; 2.Y.172.157; 2.Y.172.166;
2.Y.172.169; 2.Y.172.172; 2.Y.172.175; 2.Y.172.240; 2.Y.172.244; 2.Y.175.228;
25 2.Y.175.229; 2.Y.175.230; 2.Y.175.231; 2.Y.175.236; 2.Y.175.237; 2.Y.175.238;
2.Y.175.239; 2.Y.175.154; 2.Y.175.157; 2.Y.175.166; 2.Y.175.169; 2.Y.175.172;
2.Y.175.175; 2.Y.175.240; 2.Y.175.244; 2.Y.240.228; 2.Y.240.229; 2.Y.240.230;
2.Y.240.231; 2.Y.240.236; 2.Y.240.237; 2.Y.240.238; 2.Y.240.239; 2.Y.240.154;
2.Y.240.157; 2.Y.240.166; 2.Y.240.169; 2.Y.240.172; 2.Y.240.175; 2.Y.240.240;
30 2.Y.240.244; 2.Y.244.228; 2.Y.244.229; 2.Y.244.230; 2.Y.244.231; 2.Y.244.236;

2.Y.244.237; 2.Y.244.238; 2.Y.244.239; 2.Y.244.154; 2.Y.244.157; 2.Y.244.166;
2.Y.244.169; 2.Y.244.172; 2.Y.244.175; 2.Y.244.240; 2.Y.244.244;

Prodrugs of 3.B

- 5 3.B.228.228; 3.B.228.229; 3.B.228.230; 3.B.228.231; 3.B.228.236;
3.B.228.237; 3.B.228.238; 3.B.228.239; 3.B.228.154; 3.B.228.157; 3.B.228.166;
3.B.228.169; 3.B.228.172; 3.B.228.175; 3.B.228.240; 3.B.228.244; 3.B.229.228;
3.B.229.229; 3.B.229.230; 3.B.229.231; 3.B.229.236; 3.B.229.237; 3.B.229.238;
3.B.229.239; 3.B.229.154; 3.B.229.157; 3.B.229.166; 3.B.229.169; 3.B.229.172;
10 3.B.229.175; 3.B.229.240; 3.B.229.244; 3.B.230.228; 3.B.230.229; 3.B.230.230;
3.B.230.231; 3.B.230.236; 3.B.230.237; 3.B.230.238; 3.B.230.239; 3.B.230.154;
3.B.230.157; 3.B.230.166; 3.B.230.169; 3.B.230.172; 3.B.230.175; 3.B.230.240;
3.B.230.244; 3.B.231.228; 3.B.231.229; 3.B.231.230; 3.B.231.231; 3.B.231.236;
3.B.231.237; 3.B.231.238; 3.B.231.239; 3.B.231.154; 3.B.231.157; 3.B.231.166;
15 3.B.231.169; 3.B.231.172; 3.B.231.175; 3.B.231.240; 3.B.231.244; 3.B.236.228;
3.B.236.229; 3.B.236.230; 3.B.236.231; 3.B.236.236; 3.B.236.237; 3.B.236.238;
3.B.236.239; 3.B.236.154; 3.B.236.157; 3.B.236.166; 3.B.236.169; 3.B.236.172;
3.B.236.175; 3.B.236.240; 3.B.236.244; 3.B.237.228; 3.B.237.229; 3.B.237.230;
3.B.237.231; 3.B.237.236; 3.B.237.237; 3.B.237.238; 3.B.237.239; 3.B.237.154;
20 3.B.237.157; 3.B.237.166; 3.B.237.169; 3.B.237.172; 3.B.237.175; 3.B.237.240;
3.B.237.244; 3.B.238.228; 3.B.238.229; 3.B.238.230; 3.B.238.231; 3.B.238.236;
3.B.238.237; 3.B.238.238; 3.B.238.239; 3.B.238.154; 3.B.238.157; 3.B.238.166;
3.B.238.169; 3.B.238.172; 3.B.238.175; 3.B.238.240; 3.B.238.244; 3.B.239.228;
3.B.239.229; 3.B.239.230; 3.B.239.231; 3.B.239.236; 3.B.239.237; 3.B.239.238;
25 3.B.239.239; 3.B.239.154; 3.B.239.157; 3.B.239.166; 3.B.239.169; 3.B.239.172;
3.B.239.175; 3.B.239.240; 3.B.239.244; 3.B.154.228; 3.B.154.229; 3.B.154.230;
3.B.154.231; 3.B.154.236; 3.B.154.237; 3.B.154.238; 3.B.154.239; 3.B.154.154;
3.B.154.157; 3.B.154.166; 3.B.154.169; 3.B.154.172; 3.B.154.175; 3.B.154.240;
3.B.154.244; 3.B.157.228; 3.B.157.229; 3.B.157.230; 3.B.157.231; 3.B.157.236;
30 3.B.157.237; 3.B.157.238; 3.B.157.239; 3.B.157.154; 3.B.157.157; 3.B.157.166;

3.B.157.169; 3.B.157.172; 3.B.157.175; 3.B.157.240; 3.B.157.244; 3.B.166.228;
3.B.166.229; 3.B.166.230; 3.B.166.231; 3.B.166.236; 3.B.166.237; 3.B.166.238;
3.B.166.239; 3.B.166.154; 3.B.166.157; 3.B.166.166; 3.B.166.169; 3.B.166.172;
3.B.166.175; 3.B.166.240; 3.B.166.244; 3.B.169.228; 3.B.169.229; 3.B.169.230;
5 3.B.169.231; 3.B.169.236; 3.B.169.237; 3.B.169.238; 3.B.169.239; 3.B.169.154;
3.B.169.157; 3.B.169.166; 3.B.169.169; 3.B.169.172; 3.B.169.175; 3.B.169.240;
3.B.169.244; 3.B.172.228; 3.B.172.229; 3.B.172.230; 3.B.172.231; 3.B.172.236;
3.B.172.237; 3.B.172.238; 3.B.172.239; 3.B.172.154; 3.B.172.157; 3.B.172.166;
3.B.172.169; 3.B.172.172; 3.B.172.175; 3.B.172.240; 3.B.172.244; 3.B.175.228;
10 3.B.175.229; 3.B.175.230; 3.B.175.231; 3.B.175.236; 3.B.175.237; 3.B.175.238;
3.B.175.239; 3.B.175.154; 3.B.175.157; 3.B.175.166; 3.B.175.169; 3.B.175.172;
3.B.175.175; 3.B.175.240; 3.B.175.244; 3.B.240.228; 3.B.240.229; 3.B.240.230;
3.B.240.231; 3.B.240.236; 3.B.240.237; 3.B.240.238; 3.B.240.239; 3.B.240.154;
3.B.240.157; 3.B.240.166; 3.B.240.169; 3.B.240.172; 3.B.240.175; 3.B.240.240;
15 3.B.240.244; 3.B.244.228; 3.B.244.229; 3.B.244.230; 3.B.244.231; 3.B.244.236;
3.B.244.237; 3.B.244.238; 3.B.244.239; 3.B.244.154; 3.B.244.157; 3.B.244.166;
3.B.244.169; 3.B.244.172; 3.B.244.175; 3.B.244.240; 3.B.244.244;

Prodrugs of 3.D

20 3.D.228.228; 3.D.228.229; 3.D.228.230; 3.D.228.231; 3.D.228.236;
3.D.228.237; 3.D.228.238; 3.D.228.239; 3.D.228.154; 3.D.228.157;
3.D.228.166; 3.D.228.169; 3.D.228.172; 3.D.228.175; 3.D.228.240;
3.D.228.244; 3.D.229.228; 3.D.229.229; 3.D.229.230; 3.D.229.231;
3.D.229.236; 3.D.229.237; 3.D.229.238; 3.D.229.239; 3.D.229.154;
25 3.D.229.157; 3.D.229.166; 3.D.229.169; 3.D.229.172; 3.D.229.175;
3.D.229.240; 3.D.229.244; 3.D.230.228; 3.D.230.229; 3.D.230.230;
3.D.230.231; 3.D.230.236; 3.D.230.237; 3.D.230.238; 3.D.230.239;
3.D.230.154; 3.D.230.157; 3.D.230.166; 3.D.230.169; 3.D.230.172;
3.D.230.175; 3.D.230.240; 3.D.230.244; 3.D.231.228; 3.D.231.229;
30 3.D.231.230; 3.D.231.231; 3.D.231.236; 3.D.231.237; 3.D.231.238;

3.D.231.239; 3.D.231.154; 3.D.231.157; 3.D.231.166; 3.D.231.169;
3.D.231.172; 3.D.231.175; 3.D.231.240; 3.D.231.244; 3.D.236.228;
3.D.236.229; 3.D.236.230; 3.D.236.231; 3.D.236.236; 3.D.236.237;
3.D.236.238; 3.D.236.239; 3.D.236.154; 3.D.236.157; 3.D.236.166;
5 3.D.236.169; 3.D.236.172; 3.D.236.175; 3.D.236.240; 3.D.236.244;
3.D.237.228; 3.D.237.229; 3.D.237.230; 3.D.237.231; 3.D.237.236;
3.D.237.237; 3.D.237.238; 3.D.237.239; 3.D.237.154; 3.D.237.157;
3.D.237.166; 3.D.237.169; 3.D.237.172; 3.D.237.175; 3.D.237.240;
3.D.237.244; 3.D.238.228; 3.D.238.229; 3.D.238.230; 3.D.238.231;
10 3.D.238.236; 3.D.238.237; 3.D.238.238; 3.D.238.239; 3.D.238.154;
3.D.238.157; 3.D.238.166; 3.D.238.169; 3.D.238.172; 3.D.238.175;
3.D.238.240; 3.D.238.244; 3.D.239.228; 3.D.239.229; 3.D.239.230;
3.D.239.231; 3.D.239.236; 3.D.239.237; 3.D.239.238; 3.D.239.239;
3.D.239.154; 3.D.239.157; 3.D.239.166; 3.D.239.169; 3.D.239.172;
15 3.D.239.175; 3.D.239.240; 3.D.239.244; 3.D.154.228; 3.D.154.229;
3.D.154.230; 3.D.154.231; 3.D.154.236; 3.D.154.237; 3.D.154.238;
3.D.154.239; 3.D.154.154; 3.D.154.157; 3.D.154.166; 3.D.154.169;
3.D.154.172; 3.D.154.175; 3.D.154.240; 3.D.154.244; 3.D.157.228;
3.D.157.229; 3.D.157.230; 3.D.157.231; 3.D.157.236; 3.D.157.237;
20 3.D.157.238; 3.D.157.239; 3.D.157.154; 3.D.157.157; 3.D.157.166;
3.D.157.169; 3.D.157.172; 3.D.157.175; 3.D.157.240; 3.D.157.244;
3.D.166.228; 3.D.166.229; 3.D.166.230; 3.D.166.231; 3.D.166.236;
3.D.166.237; 3.D.166.238; 3.D.166.239; 3.D.166.154; 3.D.166.157;
3.D.166.166; 3.D.166.169; 3.D.166.172; 3.D.166.175; 3.D.166.240;
25 3.D.166.244; 3.D.169.228; 3.D.169.229; 3.D.169.230; 3.D.169.231;
3.D.169.236; 3.D.169.237; 3.D.169.238; 3.D.169.239; 3.D.169.154;
3.D.169.157; 3.D.169.166; 3.D.169.169; 3.D.169.172; 3.D.169.175;
3.D.169.240; 3.D.169.244; 3.D.172.228; 3.D.172.229; 3.D.172.230;
3.D.172.231; 3.D.172.236; 3.D.172.237; 3.D.172.238; 3.D.172.239;
30 3.D.172.154; 3.D.172.157; 3.D.172.166; 3.D.172.169; 3.D.172.172;

3.D.172.175; 3.D.172.240; 3.D.172.244; 3.D.175.228; 3.D.175.229;
3.D.175.230; 3.D.175.231; 3.D.175.236; 3.D.175.237; 3.D.175.238;
3.D.175.239; 3.D.175.154; 3.D.175.157; 3.D.175.166; 3.D.175.169;
3.D.175.172; 3.D.175.175; 3.D.175.240; 3.D.175.244; 3.D.240.228;
5 3.D.240.229; 3.D.240.230; 3.D.240.231; 3.D.240.236; 3.D.240.237;
3.D.240.238; 3.D.240.239; 3.D.240.154; 3.D.240.157; 3.D.240.166;
3.D.240.169; 3.D.240.172; 3.D.240.175; 3.D.240.240; 3.D.240.244;
3.D.244.228; 3.D.244.229; 3.D.244.230; 3.D.244.231; 3.D.244.236;
3.D.244.237; 3.D.244.238; 3.D.244.239; 3.D.244.154; 3.D.244.157;
10 3.D.244.166; 3.D.244.169; 3.D.244.172; 3.D.244.175; 3.D.244.240;
3.D.244.244;

Prodrugs of 3.E

3.E.228.228; 3.E.228.229; 3.E.228.230; 3.E.228.231; 3.E.228.236;
15 3.E.228.237; 3.E.228.238; 3.E.228.239; 3.E.228.154; 3.E.228.157; 3.E.228.166;
3.E.228.169; 3.E.228.172; 3.E.228.175; 3.E.228.240; 3.E.228.244; 3.E.229.228;
3.E.229.229; 3.E.229.230; 3.E.229.231; 3.E.229.236; 3.E.229.237; 3.E.229.238;
3.E.229.239; 3.E.229.154; 3.E.229.157; 3.E.229.166; 3.E.229.169; 3.E.229.172;
3.E.229.175; 3.E.229.240; 3.E.229.244; 3.E.230.228; 3.E.230.229; 3.E.230.230;
20 3.E.230.231; 3.E.230.236; 3.E.230.237; 3.E.230.238; 3.E.230.239; 3.E.230.154;
3.E.230.157; 3.E.230.166; 3.E.230.169; 3.E.230.172; 3.E.230.175; 3.E.230.240;
3.E.230.244; 3.E.231.228; 3.E.231.229; 3.E.231.230; 3.E.231.231; 3.E.231.236;
3.E.231.237; 3.E.231.238; 3.E.231.239; 3.E.231.154; 3.E.231.157; 3.E.231.166;
3.E.231.169; 3.E.231.172; 3.E.231.175; 3.E.231.240; 3.E.231.244; 3.E.236.228;
25 3.E.236.229; 3.E.236.230; 3.E.236.231; 3.E.236.236; 3.E.236.237; 3.E.236.238;
3.E.236.239; 3.E.236.154; 3.E.236.157; 3.E.236.166; 3.E.236.169; 3.E.236.172;
3.E.236.175; 3.E.236.240; 3.E.236.244; 3.E.237.228; 3.E.237.229; 3.E.237.230;
3.E.237.231; 3.E.237.236; 3.E.237.237; 3.E.237.238; 3.E.237.239; 3.E.237.154;
3.E.237.157; 3.E.237.166; 3.E.237.169; 3.E.237.172; 3.E.237.175; 3.E.237.240;
30 3.E.237.244; 3.E.238.228; 3.E.238.229; 3.E.238.230; 3.E.238.231; 3.E.238.236;

3.E.238.237; 3.E.238.238; 3.E.238.239; 3.E.238.154; 3.E.238.157; 3.E.238.166;
3.E.238.169; 3.E.238.172; 3.E.238.175; 3.E.238.240; 3.E.238.244; 3.E.239.228;
3.E.239.229; 3.E.239.230; 3.E.239.231; 3.E.239.236; 3.E.239.237; 3.E.239.238;
3.E.239.239; 3.E.239.154; 3.E.239.157; 3.E.239.166; 3.E.239.169; 3.E.239.172;
5 3.E.239.175; 3.E.239.240; 3.E.239.244; 3.E.154.228; 3.E.154.229; 3.E.154.230;
3.E.154.231; 3.E.154.236; 3.E.154.237; 3.E.154.238; 3.E.154.239; 3.E.154.154;
3.E.154.157; 3.E.154.166; 3.E.154.169; 3.E.154.172; 3.E.154.175; 3.E.154.240;
3.E.154.244; 3.E.157.228; 3.E.157.229; 3.E.157.230; 3.E.157.231; 3.E.157.236;
3.E.157.237; 3.E.157.238; 3.E.157.239; 3.E.157.154; 3.E.157.157; 3.E.157.166;
10 3.E.157.169; 3.E.157.172; 3.E.157.175; 3.E.157.240; 3.E.157.244; 3.E.166.228;
3.E.166.229; 3.E.166.230; 3.E.166.231; 3.E.166.236; 3.E.166.237; 3.E.166.238;
3.E.166.239; 3.E.166.154; 3.E.166.157; 3.E.166.166; 3.E.166.169; 3.E.166.172;
3.E.166.175; 3.E.166.240; 3.E.166.244; 3.E.169.228; 3.E.169.229; 3.E.169.230;
3.E.169.231; 3.E.169.236; 3.E.169.237; 3.E.169.238; 3.E.169.239; 3.E.169.154;
15 3.E.169.157; 3.E.169.166; 3.E.169.169; 3.E.169.172; 3.E.169.175; 3.E.169.240;
3.E.169.244; 3.E.172.228; 3.E.172.229; 3.E.172.230; 3.E.172.231; 3.E.172.236;
3.E.172.237; 3.E.172.238; 3.E.172.239; 3.E.172.154; 3.E.172.157; 3.E.172.166;
3.E.172.169; 3.E.172.172; 3.E.172.175; 3.E.172.240; 3.E.172.244; 3.E.175.228;
3.E.175.229; 3.E.175.230; 3.E.175.231; 3.E.175.236; 3.E.175.237; 3.E.175.238;
20 3.E.175.239; 3.E.175.154; 3.E.175.157; 3.E.175.166; 3.E.175.169; 3.E.175.172;
3.E.175.175; 3.E.175.240; 3.E.175.244; 3.E.240.228; 3.E.240.229; 3.E.240.230;
3.E.240.231; 3.E.240.236; 3.E.240.237; 3.E.240.238; 3.E.240.239; 3.E.240.154;
3.E.240.157; 3.E.240.166; 3.E.240.169; 3.E.240.172; 3.E.240.175; 3.E.240.240;
3.E.240.244; 3.E.244.228; 3.E.244.229; 3.E.244.230; 3.E.244.231; 3.E.244.236;
25 3.E.244.237; 3.E.244.238; 3.E.244.239; 3.E.244.154; 3.E.244.157; 3.E.244.166;
3.E.244.169; 3.E.244.172; 3.E.244.175; 3.E.244.240; 3.E.244.244;

Prodrugs of 3.G

3.G.228.228; 3.G.228.229; 3.G.228.230; 3.G.228.231; 3.G.228.236;
30 3.G.228.237; 3.G.228.238; 3.G.228.239; 3.G.228.154; 3.G.228.157;

3.G.228.166; 3.G.228.169; 3.G.228.172; 3.G.228.175; 3.G.228.240;
3.G.228.244; 3.G.229.228; 3.G.229.229; 3.G.229.230; 3.G.229.231;
3.G.229.236; 3.G.229.237; 3.G.229.238; 3.G.229.239; 3.G.229.154;
3.G.229.157; 3.G.229.166; 3.G.229.169; 3.G.229.172; 3.G.229.175;
5 3.G.229.240; 3.G.229.244; 3.G.230.228; 3.G.230.229; 3.G.230.230;
3.G.230.231; 3.G.230.236; 3.G.230.237; 3.G.230.238; 3.G.230.239;
3.G.230.154; 3.G.230.157; 3.G.230.166; 3.G.230.169; 3.G.230.172;
3.G.230.175; 3.G.230.240; 3.G.230.244; 3.G.231.228; 3.G.231.229;
3.G.231.230; 3.G.231.231; 3.G.231.236; 3.G.231.237; 3.G.231.238;
10 3.G.231.239; 3.G.231.154; 3.G.231.157; 3.G.231.166; 3.G.231.169;
3.G.231.172; 3.G.231.175; 3.G.231.240; 3.G.231.244; 3.G.236.228;
3.G.236.229; 3.G.236.230; 3.G.236.231; 3.G.236.236; 3.G.236.237;
3.G.236.238; 3.G.236.239; 3.G.236.154; 3.G.236.157; 3.G.236.166;
3.G.236.169; 3.G.236.172; 3.G.236.175; 3.G.236.240; 3.G.236.244;
15 3.G.237.228; 3.G.237.229; 3.G.237.230; 3.G.237.231; 3.G.237.236;
3.G.237.237; 3.G.237.238; 3.G.237.239; 3.G.237.154; 3.G.237.157;
3.G.237.166; 3.G.237.169; 3.G.237.172; 3.G.237.175; 3.G.237.240;
3.G.237.244; 3.G.238.228; 3.G.238.229; 3.G.238.230; 3.G.238.231;
3.G.238.236; 3.G.238.237; 3.G.238.238; 3.G.238.239; 3.G.238.154;
20 3.G.238.157; 3.G.238.166; 3.G.238.169; 3.G.238.172; 3.G.238.175;
3.G.238.240; 3.G.238.244; 3.G.239.228; 3.G.239.229; 3.G.239.230;
3.G.239.231; 3.G.239.236; 3.G.239.237; 3.G.239.238; 3.G.239.239;
3.G.239.154; 3.G.239.157; 3.G.239.166; 3.G.239.169; 3.G.239.172;
3.G.239.175; 3.G.239.240; 3.G.239.244; 3.G.154.228; 3.G.154.229;
25 3.G.154.230; 3.G.154.231; 3.G.154.236; 3.G.154.237; 3.G.154.238;
3.G.154.239; 3.G.154.154; 3.G.154.157; 3.G.154.166; 3.G.154.169;
3.G.154.172; 3.G.154.175; 3.G.154.240; 3.G.154.244; 3.G.157.228;
3.G.157.229; 3.G.157.230; 3.G.157.231; 3.G.157.236; 3.G.157.237;
3.G.157.238; 3.G.157.239; 3.G.157.154; 3.G.157.157; 3.G.157.166;
30 3.G.157.169; 3.G.157.172; 3.G.157.175; 3.G.157.240; 3.G.157.244;

- 3.G.166.228; 3.G.166.229; 3.G.166.230; 3.G.166.231; 3.G.166.236;
 3.G.166.237; 3.G.166.238; 3.G.166.239; 3.G.166.154; 3.G.166.157;
 3.G.166.166; 3.G.166.169; 3.G.166.172; 3.G.166.175; 3.G.166.240;
 3.G.166.244; 3.G.169.228; 3.G.169.229; 3.G.169.230; 3.G.169.231;
 5 3.G.169.236; 3.G.169.237; 3.G.169.238; 3.G.169.239; 3.G.169.154;
 3.G.169.157; 3.G.169.166; 3.G.169.169; 3.G.169.172; 3.G.169.175;
 3.G.169.240; 3.G.169.244; 3.G.172.228; 3.G.172.229; 3.G.172.230;
 3.G.172.231; 3.G.172.236; 3.G.172.237; 3.G.172.238; 3.G.172.239;
 3.G.172.154; 3.G.172.157; 3.G.172.166; 3.G.172.169; 3.G.172.172;
 10 3.G.172.175; 3.G.172.240; 3.G.172.244; 3.G.175.228; 3.G.175.229;
 3.G.175.230; 3.G.175.231; 3.G.175.236; 3.G.175.237; 3.G.175.238;
 3.G.175.239; 3.G.175.154; 3.G.175.157; 3.G.175.166; 3.G.175.169;
 3.G.175.172; 3.G.175.175; 3.G.175.240; 3.G.175.244; 3.G.240.228;
 3.G.240.229; 3.G.240.230; 3.G.240.231; 3.G.240.236; 3.G.240.237;
 15 3.G.240.238; 3.G.240.239; 3.G.240.154; 3.G.240.157; 3.G.240.166;
 3.G.240.169; 3.G.240.172; 3.G.240.175; 3.G.240.240; 3.G.240.244;
 3.G.244.228; 3.G.244.229; 3.G.244.230; 3.G.244.231; 3.G.244.236;
 3.G.244.237; 3.G.244.238; 3.G.244.239; 3.G.244.154; 3.G.244.157;
 3.G.244.166; 3.G.244.169; 3.G.244.172; 3.G.244.175; 3.G.244.240;
 20 3.G.244.244;

Prodrugs of 3.I

- 3.I.228.228; 3.I.228.229; 3.I.228.230; 3.I.228.231; 3.I.228.236; 3.I.228.237;
 3.I.228.238; 3.I.228.239; 3.I.228.154; 3.I.228.157; 3.I.228.166; 3.I.228.169;
 25 3.I.228.172; 3.I.228.175; 3.I.228.240; 3.I.228.244; 3.I.229.228; 3.I.229.229;
 3.I.229.230; 3.I.229.231; 3.I.229.236; 3.I.229.237; 3.I.229.238; 3.I.229.239;
 3.I.229.154; 3.I.229.157; 3.I.229.166; 3.I.229.169; 3.I.229.172; 3.I.229.175;
 3.I.229.240; 3.I.229.244; 3.I.230.228; 3.I.230.229; 3.I.230.230; 3.I.230.231;
 3.I.230.236; 3.I.230.237; 3.I.230.238; 3.I.230.239; 3.I.230.154; 3.I.230.157;
 30 3.I.230.166; 3.I.230.169; 3.I.230.172; 3.I.230.175; 3.I.230.240; 3.I.230.244;

- 3.I.231.228; 3.I.231.229; 3.I.231.230; 3.I.231.231; 3.I.231.236; 3.I.231.237;
3.I.231.238; 3.I.231.239; 3.I.231.154; 3.I.231.157; 3.I.231.166; 3.I.231.169;
3.I.231.172; 3.I.231.175; 3.I.231.240; 3.I.231.244; 3.I.236.228; 3.I.236.229;
3.I.236.230; 3.I.236.231; 3.I.236.236; 3.I.236.237; 3.I.236.238; 3.I.236.239;
5 3.I.236.154; 3.I.236.157; 3.I.236.166; 3.I.236.169; 3.I.236.172; 3.I.236.175;
3.I.236.240; 3.I.236.244; 3.I.237.228; 3.I.237.229; 3.I.237.230; 3.I.237.231;
3.I.237.236; 3.I.237.237; 3.I.237.238; 3.I.237.239; 3.I.237.154; 3.I.237.157;
3.I.237.166; 3.I.237.169; 3.I.237.172; 3.I.237.175; 3.I.237.240; 3.I.237.244;
3.I.238.228; 3.I.238.229; 3.I.238.230; 3.I.238.231; 3.I.238.236; 3.I.238.237;
10 3.I.238.238; 3.I.238.239; 3.I.238.154; 3.I.238.157; 3.I.238.166; 3.I.238.169;
3.I.238.172; 3.I.238.175; 3.I.238.240; 3.I.238.244; 3.I.239.228; 3.I.239.229;
3.I.239.230; 3.I.239.231; 3.I.239.236; 3.I.239.237; 3.I.239.238; 3.I.239.239;
3.I.239.154; 3.I.239.157; 3.I.239.166; 3.I.239.169; 3.I.239.172; 3.I.239.175;
3.I.239.240; 3.I.239.244; 3.I.154.228; 3.I.154.229; 3.I.154.230; 3.I.154.231;
15 3.I.154.236; 3.I.154.237; 3.I.154.238; 3.I.154.239; 3.I.154.154; 3.I.154.157;
3.I.154.166; 3.I.154.169; 3.I.154.172; 3.I.154.175; 3.I.154.240; 3.I.154.244;
3.I.157.228; 3.I.157.229; 3.I.157.230; 3.I.157.231; 3.I.157.236; 3.I.157.237;
3.I.157.238; 3.I.157.239; 3.I.157.154; 3.I.157.157; 3.I.157.166; 3.I.157.169;
3.I.157.172; 3.I.157.175; 3.I.157.240; 3.I.157.244; 3.I.166.228; 3.I.166.229;
20 3.I.166.230; 3.I.166.231; 3.I.166.236; 3.I.166.237; 3.I.166.238; 3.I.166.239;
3.I.166.154; 3.I.166.157; 3.I.166.166; 3.I.166.169; 3.I.166.172; 3.I.166.175;
3.I.166.240; 3.I.166.244; 3.I.169.228; 3.I.169.229; 3.I.169.230; 3.I.169.231;
3.I.169.236; 3.I.169.237; 3.I.169.238; 3.I.169.239; 3.I.169.154; 3.I.169.157;
3.I.169.166; 3.I.169.169; 3.I.169.172; 3.I.169.175; 3.I.169.240; 3.I.169.244;
25 3.I.172.228; 3.I.172.229; 3.I.172.230; 3.I.172.231; 3.I.172.236; 3.I.172.237;
3.I.172.238; 3.I.172.239; 3.I.172.154; 3.I.172.157; 3.I.172.166; 3.I.172.169;
3.I.172.172; 3.I.172.175; 3.I.172.240; 3.I.172.244; 3.I.175.228; 3.I.175.229;
3.I.175.230; 3.I.175.231; 3.I.175.236; 3.I.175.237; 3.I.175.238; 3.I.175.239;
3.I.175.154; 3.I.175.157; 3.I.175.166; 3.I.175.169; 3.I.175.172; 3.I.175.175;
30 3.I.175.240; 3.I.175.244; 3.I.240.228; 3.I.240.229; 3.I.240.230; 3.I.240.231;

3.I.240.236; 3.I.240.237; 3.I.240.238; 3.I.240.239; 3.I.240.154; 3.I.240.157;
3.I.240.166; 3.I.240.169; 3.I.240.172; 3.I.240.175; 3.I.240.240; 3.I.240.244;
3.I.244.228; 3.I.244.229; 3.I.244.230; 3.I.244.231; 3.I.244.236; 3.I.244.237;
3.I.244.238; 3.I.244.239; 3.I.244.154; 3.I.244.157; 3.I.244.166; 3.I.244.169;
5 3.I.244.172; 3.I.244.175; 3.I.244.240; 3.I.244.244;

Prodrugs of 3.I

3.J.228.228; 3.J.228.229; 3.J.228.230; 3.J.228.231; 3.J.228.236; 3.J.228.237;
3.J.228.238; 3.J.228.239; 3.J.228.154; 3.J.228.157; 3.J.228.166; 3.J.228.169;
10 3.J.228.172; 3.J.228.175; 3.J.228.240; 3.J.228.244; 3.J.229.228; 3.J.229.229;
3.J.229.230; 3.J.229.231; 3.J.229.236; 3.J.229.237; 3.J.229.238; 3.J.229.239;
3.J.229.154; 3.J.229.157; 3.J.229.166; 3.J.229.169; 3.J.229.172; 3.J.229.175;
3.J.229.240; 3.J.229.244; 3.J.230.228; 3.J.230.229; 3.J.230.230; 3.J.230.231;
3.J.230.236; 3.J.230.237; 3.J.230.238; 3.J.230.239; 3.J.230.154; 3.J.230.157;
15 3.J.230.166; 3.J.230.169; 3.J.230.172; 3.J.230.175; 3.J.230.240; 3.J.230.244;
3.J.231.228; 3.J.231.229; 3.J.231.230; 3.J.231.231; 3.J.231.236; 3.J.231.237;
3.J.231.238; 3.J.231.239; 3.J.231.154; 3.J.231.157; 3.J.231.166; 3.J.231.169;
3.J.231.172; 3.J.231.175; 3.J.231.240; 3.J.231.244; 3.J.236.228; 3.J.236.229;
3.J.236.230; 3.J.236.231; 3.J.236.236; 3.J.236.237; 3.J.236.238; 3.J.236.239;
20 3.J.236.154; 3.J.236.157; 3.J.236.166; 3.J.236.169; 3.J.236.172; 3.J.236.175;
3.J.236.240; 3.J.236.244; 3.J.237.228; 3.J.237.229; 3.J.237.230; 3.J.237.231;
3.J.237.236; 3.J.237.237; 3.J.237.238; 3.J.237.239; 3.J.237.154; 3.J.237.157;
3.J.237.166; 3.J.237.169; 3.J.237.172; 3.J.237.175; 3.J.237.240; 3.J.237.244;
3.J.238.228; 3.J.238.229; 3.J.238.230; 3.J.238.231; 3.J.238.236; 3.J.238.237;
25 3.J.238.238; 3.J.238.239; 3.J.238.154; 3.J.238.157; 3.J.238.166; 3.J.238.169;
3.J.238.172; 3.J.238.175; 3.J.238.240; 3.J.238.244; 3.J.239.228; 3.J.239.229;
3.J.239.230; 3.J.239.231; 3.J.239.236; 3.J.239.237; 3.J.239.238; 3.J.239.239;
3.J.239.154; 3.J.239.157; 3.J.239.166; 3.J.239.169; 3.J.239.172; 3.J.239.175;
3.J.239.240; 3.J.239.244; 3.J.154.228; 3.J.154.229; 3.J.154.230; 3.J.154.231;
30 3.J.154.236; 3.J.154.237; 3.J.154.238; 3.J.154.239; 3.J.154.154; 3.J.154.157;

3.J.154.166; 3.J.154.169; 3.J.154.172; 3.J.154.175; 3.J.154.240; 3.J.154.244;
3.J.157.228; 3.J.157.229; 3.J.157.230; 3.J.157.231; 3.J.157.236; 3.J.157.237;
3.J.157.238; 3.J.157.239; 3.J.157.154; 3.J.157.157; 3.J.157.166; 3.J.157.169;
3.J.157.172; 3.J.157.175; 3.J.157.240; 3.J.157.244; 3.J.166.228; 3.J.166.229;
5 3.J.166.230; 3.J.166.231; 3.J.166.236; 3.J.166.237; 3.J.166.238; 3.J.166.239;
3.J.166.154; 3.J.166.157; 3.J.166.166; 3.J.166.169; 3.J.166.172; 3.J.166.175;
3.J.166.240; 3.J.166.244; 3.J.169.228; 3.J.169.229; 3.J.169.230; 3.J.169.231;
3.J.169.236; 3.J.169.237; 3.J.169.238; 3.J.169.239; 3.J.169.154; 3.J.169.157;
3.J.169.166; 3.J.169.169; 3.J.169.172; 3.J.169.175; 3.J.169.240; 3.J.169.244;
10 3.J.172.228; 3.J.172.229; 3.J.172.230; 3.J.172.231; 3.J.172.236; 3.J.172.237;
3.J.172.238; 3.J.172.239; 3.J.172.154; 3.J.172.157; 3.J.172.166; 3.J.172.169;
3.J.172.172; 3.J.172.175; 3.J.172.240; 3.J.172.244; 3.J.175.228; 3.J.175.229;
3.J.175.230; 3.J.175.231; 3.J.175.236; 3.J.175.237; 3.J.175.238; 3.J.175.239;
3.J.175.154; 3.J.175.157; 3.J.175.166; 3.J.175.169; 3.J.175.172; 3.J.175.175;
15 3.J.175.240; 3.J.175.244; 3.J.240.228; 3.J.240.229; 3.J.240.230; 3.J.240.231;
3.J.240.236; 3.J.240.237; 3.J.240.238; 3.J.240.239; 3.J.240.154; 3.J.240.157;
3.J.240.166; 3.J.240.169; 3.J.240.172; 3.J.240.175; 3.J.240.240; 3.J.240.244;
3.J.244.228; 3.J.244.229; 3.J.244.230; 3.J.244.231; 3.J.244.236; 3.J.244.237;
3.J.244.238; 3.J.244.239; 3.J.244.154; 3.J.244.157; 3.J.244.166; 3.J.244.169;
20 3.J.244.172; 3.J.244.175; 3.J.244.240; 3.J.244.244;

Prodrugs of 3.L

3.L.228.228; 3.L.228.229; 3.L.228.230; 3.L.228.231; 3.L.228.236;
3.L.228.237; 3.L.228.238; 3.L.228.239; 3.L.228.154; 3.L.228.157; 3.L.228.166;
25 3.L.228.169; 3.L.228.172; 3.L.228.175; 3.L.228.240; 3.L.228.244; 3.L.229.228;
3.L.229.229; 3.L.229.230; 3.L.229.231; 3.L.229.236; 3.L.229.237; 3.L.229.238;
3.L.229.239; 3.L.229.154; 3.L.229.157; 3.L.229.166; 3.L.229.169; 3.L.229.172;
3.L.229.175; 3.L.229.240; 3.L.229.244; 3.L.230.228; 3.L.230.229; 3.L.230.230;
3.L.230.231; 3.L.230.236; 3.L.230.237; 3.L.230.238; 3.L.230.239; 3.L.230.154;
30 3.L.230.157; 3.L.230.166; 3.L.230.169; 3.L.230.172; 3.L.230.175; 3.L.230.240;

3.L.230.244; 3.L.231.228; 3.L.231.229; 3.L.231.230; 3.L.231.231; 3.L.231.236;
3.L.231.237; 3.L.231.238; 3.L.231.239; 3.L.231.154; 3.L.231.157; 3.L.231.166;
3.L.231.169; 3.L.231.172; 3.L.231.175; 3.L.231.240; 3.L.231.244; 3.L.236.228;
3.L.236.229; 3.L.236.230; 3.L.236.231; 3.L.236.236; 3.L.236.237; 3.L.236.238;
5 3.L.236.239; 3.L.236.154; 3.L.236.157; 3.L.236.166; 3.L.236.169; 3.L.236.172;
3.L.236.175; 3.L.236.240; 3.L.236.244; 3.L.237.228; 3.L.237.229; 3.L.237.230;
3.L.237.231; 3.L.237.236; 3.L.237.237; 3.L.237.238; 3.L.237.239; 3.L.237.154;
3.L.237.157; 3.L.237.166; 3.L.237.169; 3.L.237.172; 3.L.237.175; 3.L.237.240;
3.L.237.244; 3.L.238.228; 3.L.238.229; 3.L.238.230; 3.L.238.231; 3.L.238.236;
10 3.L.238.237; 3.L.238.238; 3.L.238.239; 3.L.238.154; 3.L.238.157; 3.L.238.166;
3.L.238.169; 3.L.238.172; 3.L.238.175; 3.L.238.240; 3.L.238.244; 3.L.239.228;
3.L.239.229; 3.L.239.230; 3.L.239.231; 3.L.239.236; 3.L.239.237; 3.L.239.238;
3.L.239.239; 3.L.239.154; 3.L.239.157; 3.L.239.166; 3.L.239.169; 3.L.239.172;
3.L.239.175; 3.L.239.240; 3.L.239.244; 3.L.154.228; 3.L.154.229; 3.L.154.230;
15 3.L.154.231; 3.L.154.236; 3.L.154.237; 3.L.154.238; 3.L.154.239; 3.L.154.154;
3.L.154.157; 3.L.154.166; 3.L.154.169; 3.L.154.172; 3.L.154.175; 3.L.154.240;
3.L.154.244; 3.L.157.228; 3.L.157.229; 3.L.157.230; 3.L.157.231; 3.L.157.236;
3.L.157.237; 3.L.157.238; 3.L.157.239; 3.L.157.154; 3.L.157.157; 3.L.157.166;
3.L.157.169; 3.L.157.172; 3.L.157.175; 3.L.157.240; 3.L.157.244; 3.L.166.228;
20 3.L.166.229; 3.L.166.230; 3.L.166.231; 3.L.166.236; 3.L.166.237; 3.L.166.238;
3.L.166.239; 3.L.166.154; 3.L.166.157; 3.L.166.166; 3.L.166.169; 3.L.166.172;
3.L.166.175; 3.L.166.240; 3.L.166.244; 3.L.169.228; 3.L.169.229; 3.L.169.230;
3.L.169.231; 3.L.169.236; 3.L.169.237; 3.L.169.238; 3.L.169.239; 3.L.169.154;
3.L.169.157; 3.L.169.166; 3.L.169.169; 3.L.169.172; 3.L.169.175; 3.L.169.240;
25 3.L.169.244; 3.L.172.228; 3.L.172.229; 3.L.172.230; 3.L.172.231; 3.L.172.236;
3.L.172.237; 3.L.172.238; 3.L.172.239; 3.L.172.154; 3.L.172.157; 3.L.172.166;
3.L.172.169; 3.L.172.172; 3.L.172.175; 3.L.172.240; 3.L.172.244; 3.L.175.228;
3.L.175.229; 3.L.175.230; 3.L.175.231; 3.L.175.236; 3.L.175.237; 3.L.175.238;
3.L.175.239; 3.L.175.154; 3.L.175.157; 3.L.175.166; 3.L.175.169; 3.L.175.172;
30 3.L.175.175; 3.L.175.240; 3.L.175.244; 3.L.240.228; 3.L.240.229; 3.L.240.230;

3.L.240.231; 3.L.240.236; 3.L.240.237; 3.L.240.238; 3.L.240.239; 3.L.240.154;
3.L.240.157; 3.L.240.166; 3.L.240.169; 3.L.240.172; 3.L.240.175; 3.L.240.240;
3.L.240.244; 3.L.244.228; 3.L.244.229; 3.L.244.230; 3.L.244.231; 3.L.244.236;
3.L.244.237; 3.L.244.238; 3.L.244.239; 3.L.244.154; 3.L.244.157; 3.L.244.166;
5 3.L.244.169; 3.L.244.172; 3.L.244.175; 3.L.244.240; 3.L.244.244;

Prodrugs of 3.O

3.O.228.228; 3.O.228.229; 3.O.228.230; 3.O.228.231; 3.O.228.236;
3.O.228.237; 3.O.228.238; 3.O.228.239; 3.O.228.154; 3.O.228.157;
10 3.O.228.166; 3.O.228.169; 3.O.228.172; 3.O.228.175; 3.O.228.240;
3.O.228.244; 3.O.229.228; 3.O.229.229; 3.O.229.230; 3.O.229.231;
3.O.229.236; 3.O.229.237; 3.O.229.238; 3.O.229.239; 3.O.229.154;
3.O.229.157; 3.O.229.166; 3.O.229.169; 3.O.229.172; 3.O.229.175;
3.O.229.240; 3.O.229.244; 3.O.230.228; 3.O.230.229; 3.O.230.230;
15 3.O.230.231; 3.O.230.236; 3.O.230.237; 3.O.230.238; 3.O.230.239;
3.O.230.154; 3.O.230.157; 3.O.230.166; 3.O.230.169; 3.O.230.172;
3.O.230.175; 3.O.230.240; 3.O.230.244; 3.O.231.228; 3.O.231.229;
3.O.231.230; 3.O.231.231; 3.O.231.236; 3.O.231.237; 3.O.231.238;
3.O.231.239; 3.O.231.154; 3.O.231.157; 3.O.231.166; 3.O.231.169;
20 3.O.231.172; 3.O.231.175; 3.O.231.240; 3.O.231.244; 3.O.236.228;
3.O.236.229; 3.O.236.230; 3.O.236.231; 3.O.236.236; 3.O.236.237;
3.O.236.238; 3.O.236.239; 3.O.236.154; 3.O.236.157; 3.O.236.166;
3.O.236.169; 3.O.236.172; 3.O.236.175; 3.O.236.240; 3.O.236.244;
3.O.237.228; 3.O.237.229; 3.O.237.230; 3.O.237.231; 3.O.237.236;
25 3.O.237.237; 3.O.237.238; 3.O.237.239; 3.O.237.154; 3.O.237.157;
3.O.237.166; 3.O.237.169; 3.O.237.172; 3.O.237.175; 3.O.237.240;
3.O.237.244; 3.O.238.228; 3.O.238.229; 3.O.238.230; 3.O.238.231;
3.O.238.236; 3.O.238.237; 3.O.238.238; 3.O.238.239; 3.O.238.154;
3.O.238.157; 3.O.238.166; 3.O.238.169; 3.O.238.172; 3.O.238.175;
30 3.O.238.240; 3.O.238.244; 3.O.239.228; 3.O.239.229; 3.O.239.230;

3.O.239.231; 3.O.239.236; 3.O.239.237; 3.O.239.238; 3.O.239.239;
3.O.239.154; 3.O.239.157; 3.O.239.166; 3.O.239.169; 3.O.239.172;
3.O.239.175; 3.O.239.240; 3.O.239.244; 3.O.154.228; 3.O.154.229;
3.O.154.230; 3.O.154.231; 3.O.154.236; 3.O.154.237; 3.O.154.238;
5 3.O.154.239; 3.O.154.154; 3.O.154.157; 3.O.154.166; 3.O.154.169;
3.O.154.172; 3.O.154.175; 3.O.154.240; 3.O.154.244; 3.O.157.228;
3.O.157.229; 3.O.157.230; 3.O.157.231; 3.O.157.236; 3.O.157.237;
3.O.157.238; 3.O.157.239; 3.O.157.154; 3.O.157.157; 3.O.157.166;
3.O.157.169; 3.O.157.172; 3.O.157.175; 3.O.157.240; 3.O.157.244;
10 3.O.166.228; 3.O.166.229; 3.O.166.230; 3.O.166.231; 3.O.166.236;
3.O.166.237; 3.O.166.238; 3.O.166.239; 3.O.166.154; 3.O.166.157;
3.O.166.166; 3.O.166.169; 3.O.166.172; 3.O.166.175; 3.O.166.240;
3.O.166.244; 3.O.169.228; 3.O.169.229; 3.O.169.230; 3.O.169.231;
3.O.169.236; 3.O.169.237; 3.O.169.238; 3.O.169.239; 3.O.169.154;
15 3.O.169.157; 3.O.169.166; 3.O.169.169; 3.O.169.172; 3.O.169.175;
3.O.169.240; 3.O.169.244; 3.O.172.228; 3.O.172.229; 3.O.172.230;
3.O.172.231; 3.O.172.236; 3.O.172.237; 3.O.172.238; 3.O.172.239;
3.O.172.154; 3.O.172.157; 3.O.172.166; 3.O.172.169; 3.O.172.172;
3.O.172.175; 3.O.172.240; 3.O.172.244; 3.O.175.228; 3.O.175.229;
20 3.O.175.230; 3.O.175.231; 3.O.175.236; 3.O.175.237; 3.O.175.238;
3.O.175.239; 3.O.175.154; 3.O.175.157; 3.O.175.166; 3.O.175.169;
3.O.175.172; 3.O.175.175; 3.O.175.240; 3.O.175.244; 3.O.240.228;
3.O.240.229; 3.O.240.230; 3.O.240.231; 3.O.240.236; 3.O.240.237;
3.O.240.238; 3.O.240.239; 3.O.240.154; 3.O.240.157; 3.O.240.166;
25 3.O.240.169; 3.O.240.172; 3.O.240.175; 3.O.240.240; 3.O.240.244;
3.O.244.228; 3.O.244.229; 3.O.244.230; 3.O.244.231; 3.O.244.236;
3.O.244.237; 3.O.244.238; 3.O.244.239; 3.O.244.154; 3.O.244.157;
3.O.244.166; 3.O.244.169; 3.O.244.172; 3.O.244.175; 3.O.244.240;
3.O.244.244;

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Prodrugs of 3.P

- 3.P.228.228; 3.P.228.229; 3.P.228.230; 3.P.228.231; 3.P.228.236;
3.P.228.237; 3.P.228.238; 3.P.228.239; 3.P.228.154; 3.P.228.157; 3.P.228.166;
3.P.228.169; 3.P.228.172; 3.P.228.175; 3.P.228.240; 3.P.228.244; 3.P.229.228;
5 3.P.229.229; 3.P.229.230; 3.P.229.231; 3.P.229.236; 3.P.229.237; 3.P.229.238;
3.P.229.239; 3.P.229.154; 3.P.229.157; 3.P.229.166; 3.P.229.169; 3.P.229.172;
3.P.229.175; 3.P.229.240; 3.P.229.244; 3.P.230.228; 3.P.230.229; 3.P.230.230;
3.P.230.231; 3.P.230.236; 3.P.230.237; 3.P.230.238; 3.P.230.239; 3.P.230.154;
3.P.230.157; 3.P.230.166; 3.P.230.169; 3.P.230.172; 3.P.230.175; 3.P.230.240;
10 3.P.230.244; 3.P.231.228; 3.P.231.229; 3.P.231.230; 3.P.231.231; 3.P.231.236;
3.P.231.237; 3.P.231.238; 3.P.231.239; 3.P.231.154; 3.P.231.157; 3.P.231.166;
3.P.231.169; 3.P.231.172; 3.P.231.175; 3.P.231.240; 3.P.231.244; 3.P.236.228;
3.P.236.229; 3.P.236.230; 3.P.236.231; 3.P.236.236; 3.P.236.237; 3.P.236.238;
3.P.236.239; 3.P.236.154; 3.P.236.157; 3.P.236.166; 3.P.236.169; 3.P.236.172;
15 3.P.236.175; 3.P.236.240; 3.P.236.244; 3.P.237.228; 3.P.237.229; 3.P.237.230;
3.P.237.231; 3.P.237.236; 3.P.237.237; 3.P.237.238; 3.P.237.239; 3.P.237.154;
3.P.237.157; 3.P.237.166; 3.P.237.169; 3.P.237.172; 3.P.237.175; 3.P.237.240;
3.P.237.244; 3.P.238.228; 3.P.238.229; 3.P.238.230; 3.P.238.231; 3.P.238.236;
3.P.238.237; 3.P.238.238; 3.P.238.239; 3.P.238.154; 3.P.238.157; 3.P.238.166;
20 3.P.238.169; 3.P.238.172; 3.P.238.175; 3.P.238.240; 3.P.238.244; 3.P.239.228;
3.P.239.229; 3.P.239.230; 3.P.239.231; 3.P.239.236; 3.P.239.237; 3.P.239.238;
3.P.239.239; 3.P.239.154; 3.P.239.157; 3.P.239.166; 3.P.239.169; 3.P.239.172;
3.P.239.175; 3.P.239.240; 3.P.239.244; 3.P.154.228; 3.P.154.229; 3.P.154.230;
3.P.154.231; 3.P.154.236; 3.P.154.237; 3.P.154.238; 3.P.154.239; 3.P.154.154;
25 3.P.154.157; 3.P.154.166; 3.P.154.169; 3.P.154.172; 3.P.154.175; 3.P.154.240;
3.P.154.244; 3.P.157.228; 3.P.157.229; 3.P.157.230; 3.P.157.231; 3.P.157.236;
3.P.157.237; 3.P.157.238; 3.P.157.239; 3.P.157.154; 3.P.157.157; 3.P.157.166;
3.P.157.169; 3.P.157.172; 3.P.157.175; 3.P.157.240; 3.P.157.244; 3.P.166.228;
3.P.166.229; 3.P.166.230; 3.P.166.231; 3.P.166.236; 3.P.166.237; 3.P.166.238;
30 3.P.166.239; 3.P.166.154; 3.P.166.157; 3.P.166.166; 3.P.166.169; 3.P.166.172;

3.P.166.175; 3.P.166.240; 3.P.166.244; 3.P.169.228; 3.P.169.229; 3.P.169.230;
 3.P.169.231; 3.P.169.236; 3.P.169.237; 3.P.169.238; 3.P.169.239; 3.P.169.154;
 3.P.169.157; 3.P.169.166; 3.P.169.169; 3.P.169.172; 3.P.169.175; 3.P.169.240;
 3.P.169.244; 3.P.172.228; 3.P.172.229; 3.P.172.230; 3.P.172.231; 3.P.172.236;
 5 3.P.172.237; 3.P.172.238; 3.P.172.239; 3.P.172.154; 3.P.172.157; 3.P.172.166;
 3.P.172.169; 3.P.172.172; 3.P.172.175; 3.P.172.240; 3.P.172.244; 3.P.175.228;
 3.P.175.229; 3.P.175.230; 3.P.175.231; 3.P.175.236; 3.P.175.237; 3.P.175.238;
 3.P.175.239; 3.P.175.154; 3.P.175.157; 3.P.175.166; 3.P.175.169; 3.P.175.172;
 3.P.175.175; 3.P.175.240; 3.P.175.244; 3.P.240.228; 3.P.240.229; 3.P.240.230;
 10 3.P.240.231; 3.P.240.236; 3.P.240.237; 3.P.240.238; 3.P.240.239; 3.P.240.154;
 3.P.240.157; 3.P.240.166; 3.P.240.169; 3.P.240.172; 3.P.240.175; 3.P.240.240;
 3.P.240.244; 3.P.244.228; 3.P.244.229; 3.P.244.230; 3.P.244.231; 3.P.244.236;
 3.P.244.237; 3.P.244.238; 3.P.244.239; 3.P.244.154; 3.P.244.157; 3.P.244.166;
 3.P.244.169; 3.P.244.172; 3.P.244.175; 3.P.244.240; 3.P.244.244;

15

Prodrugs of 3.U

3.U.228.228; 3.U.228.229; 3.U.228.230; 3.U.228.231; 3.U.228.236;
 3.U.228.237; 3.U.228.238; 3.U.228.239; 3.U.228.154; 3.U.228.157;
 3.U.228.166; 3.U.228.169; 3.U.228.172; 3.U.228.175; 3.U.228.240;
 20 3.U.228.244; 3.U.229.228; 3.U.229.229; 3.U.229.230; 3.U.229.231;
 3.U.229.236; 3.U.229.237; 3.U.229.238; 3.U.229.239; 3.U.229.154;
 3.U.229.157; 3.U.229.166; 3.U.229.169; 3.U.229.172; 3.U.229.175;
 3.U.229.240; 3.U.229.244; 3.U.230.228; 3.U.230.229; 3.U.230.230;
 3.U.230.231; 3.U.230.236; 3.U.230.237; 3.U.230.238; 3.U.230.239;
 25 3.U.230.154; 3.U.230.157; 3.U.230.166; 3.U.230.169; 3.U.230.172;
 3.U.230.175; 3.U.230.240; 3.U.230.244; 3.U.231.228; 3.U.231.229;
 3.U.231.230; 3.U.231.231; 3.U.231.236; 3.U.231.237; 3.U.231.238;
 3.U.231.239; 3.U.231.154; 3.U.231.157; 3.U.231.166; 3.U.231.169;
 3.U.231.172; 3.U.231.175; 3.U.231.240; 3.U.231.244; 3.U.236.228;
 30 3.U.236.229; 3.U.236.230; 3.U.236.231; 3.U.236.236; 3.U.236.237;

3.U.236.238; 3.U.236.239; 3.U.236.154; 3.U.236.157; 3.U.236.166;
3.U.236.169; 3.U.236.172; 3.U.236.175; 3.U.236.240; 3.U.236.244;
3.U.237.228; 3.U.237.229; 3.U.237.230; 3.U.237.231; 3.U.237.236;
3.U.237.237; 3.U.237.238; 3.U.237.239; 3.U.237.154; 3.U.237.157;
5 3.U.237.166; 3.U.237.169; 3.U.237.172; 3.U.237.175; 3.U.237.240;
3.U.237.244; 3.U.238.228; 3.U.238.229; 3.U.238.230; 3.U.238.231;
3.U.238.236; 3.U.238.237; 3.U.238.238; 3.U.238.239; 3.U.238.154;
3.U.238.157; 3.U.238.166; 3.U.238.169; 3.U.238.172; 3.U.238.175;
3.U.238.240; 3.U.238.244; 3.U.239.228; 3.U.239.229; 3.U.239.230;
10 3.U.239.231; 3.U.239.236; 3.U.239.237; 3.U.239.238; 3.U.239.239;
3.U.239.154; 3.U.239.157; 3.U.239.166; 3.U.239.169; 3.U.239.172;
3.U.239.175; 3.U.239.240; 3.U.239.244; 3.U.154.228; 3.U.154.229;
3.U.154.230; 3.U.154.231; 3.U.154.236; 3.U.154.237; 3.U.154.238;
3.U.154.239; 3.U.154.154; 3.U.154.157; 3.U.154.166; 3.U.154.169;
15 3.U.154.172; 3.U.154.175; 3.U.154.240; 3.U.154.244; 3.U.157.228;
3.U.157.229; 3.U.157.230; 3.U.157.231; 3.U.157.236; 3.U.157.237;
3.U.157.238; 3.U.157.239; 3.U.157.154; 3.U.157.157; 3.U.157.166;
3.U.157.169; 3.U.157.172; 3.U.157.175; 3.U.157.240; 3.U.157.244;
3.U.166.228; 3.U.166.229; 3.U.166.230; 3.U.166.231; 3.U.166.236;
20 3.U.166.237; 3.U.166.238; 3.U.166.239; 3.U.166.154; 3.U.166.157;
3.U.166.166; 3.U.166.169; 3.U.166.172; 3.U.166.175; 3.U.166.240;
3.U.166.244; 3.U.169.228; 3.U.169.229; 3.U.169.230; 3.U.169.231;
3.U.169.236; 3.U.169.237; 3.U.169.238; 3.U.169.239; 3.U.169.154;
3.U.169.157; 3.U.169.166; 3.U.169.169; 3.U.169.172; 3.U.169.175;
25 3.U.169.240; 3.U.169.244; 3.U.172.228; 3.U.172.229; 3.U.172.230;
3.U.172.231; 3.U.172.236; 3.U.172.237; 3.U.172.238; 3.U.172.239;
3.U.172.154; 3.U.172.157; 3.U.172.166; 3.U.172.169; 3.U.172.172;
3.U.172.175; 3.U.172.240; 3.U.172.244; 3.U.175.228; 3.U.175.229;
3.U.175.230; 3.U.175.231; 3.U.175.236; 3.U.175.237; 3.U.175.238;
30 3.U.175.239; 3.U.175.154; 3.U.175.157; 3.U.175.166; 3.U.175.169;

3.U.175.172; 3.U.175.175; 3.U.175.240; 3.U.175.244; 3.U.240.228;
3.U.240.229; 3.U.240.230; 3.U.240.231; 3.U.240.236; 3.U.240.237;
3.U.240.238; 3.U.240.239; 3.U.240.154; 3.U.240.157; 3.U.240.166;
3.U.240.169; 3.U.240.172; 3.U.240.175; 3.U.240.240; 3.U.240.244;
5 3.U.244.228; 3.U.244.229; 3.U.244.230; 3.U.244.231; 3.U.244.236;
3.U.244.237; 3.U.244.238; 3.U.244.239; 3.U.244.154; 3.U.244.157;
3.U.244.166; 3.U.244.169; 3.U.244.172; 3.U.244.175; 3.U.244.240;
3.U.244.244;

10 Prodrugs of 3.W

3.W.228.228; 3.W.228.229; 3.W.228.230; 3.W.228.231; 3.W.228.236;
3.W.228.237; 3.W.228.238; 3.W.228.239; 3.W.228.154; 3.W.228.157;
3.W.228.166; 3.W.228.169; 3.W.228.172; 3.W.228.175; 3.W.228.240;
3.W.228.244; 3.W.229.228; 3.W.229.229; 3.W.229.230; 3.W.229.231;
15 3.W.229.236; 3.W.229.237; 3.W.229.238; 3.W.229.239; 3.W.229.154;
3.W.229.157; 3.W.229.166; 3.W.229.169; 3.W.229.172; 3.W.229.175;
3.W.229.240; 3.W.229.244; 3.W.230.228; 3.W.230.229; 3.W.230.230;
3.W.230.231; 3.W.230.236; 3.W.230.237; 3.W.230.238; 3.W.230.239;
3.W.230.154; 3.W.230.157; 3.W.230.166; 3.W.230.169; 3.W.230.172;
20 3.W.230.175; 3.W.230.240; 3.W.230.244; 3.W.231.228; 3.W.231.229;
3.W.231.230; 3.W.231.231; 3.W.231.236; 3.W.231.237; 3.W.231.238;
3.W.231.239; 3.W.231.154; 3.W.231.157; 3.W.231.166; 3.W.231.169;
3.W.231.172; 3.W.231.175; 3.W.231.240; 3.W.231.244; 3.W.236.228;
3.W.236.229; 3.W.236.230; 3.W.236.231; 3.W.236.236; 3.W.236.237;
25 3.W.236.238; 3.W.236.239; 3.W.236.154; 3.W.236.157; 3.W.236.166;
3.W.236.169; 3.W.236.172; 3.W.236.175; 3.W.236.240; 3.W.236.244;
3.W.237.228; 3.W.237.229; 3.W.237.230; 3.W.237.231; 3.W.237.236;
3.W.237.237; 3.W.237.238; 3.W.237.239; 3.W.237.154; 3.W.237.157;
3.W.237.166; 3.W.237.169; 3.W.237.172; 3.W.237.175; 3.W.237.240;
30 3.W.237.244; 3.W.238.228; 3.W.238.229; 3.W.238.230; 3.W.238.231;

3.W.238.236; 3.W.238.237; 3.W.238.238; 3.W.238.239; 3.W.238.154;
3.W.238.157; 3.W.238.166; 3.W.238.169; 3.W.238.172; 3.W.238.175;
3.W.238.240; 3.W.238.244; 3.W.239.228; 3.W.239.229; 3.W.239.230;
3.W.239.231; 3.W.239.236; 3.W.239.237; 3.W.239.238; 3.W.239.239;
5 3.W.239.154; 3.W.239.157; 3.W.239.166; 3.W.239.169; 3.W.239.172;
3.W.239.175; 3.W.239.240; 3.W.239.244; 3.W.154.228; 3.W.154.229;
3.W.154.230; 3.W.154.231; 3.W.154.236; 3.W.154.237; 3.W.154.238;
3.W.154.239; 3.W.154.154; 3.W.154.157; 3.W.154.166; 3.W.154.169;
3.W.154.172; 3.W.154.175; 3.W.154.240; 3.W.154.244; 3.W.157.228;
10 3.W.157.229; 3.W.157.230; 3.W.157.231; 3.W.157.236; 3.W.157.237;
3.W.157.238; 3.W.157.239; 3.W.157.154; 3.W.157.157; 3.W.157.166;
3.W.157.169; 3.W.157.172; 3.W.157.175; 3.W.157.240; 3.W.157.244;
3.W.166.228; 3.W.166.229; 3.W.166.230; 3.W.166.231; 3.W.166.236;
3.W.166.237; 3.W.166.238; 3.W.166.239; 3.W.166.154; 3.W.166.157;
15 3.W.166.166; 3.W.166.169; 3.W.166.172; 3.W.166.175; 3.W.166.240;
3.W.166.244; 3.W.169.228; 3.W.169.229; 3.W.169.230; 3.W.169.231;
3.W.169.236; 3.W.169.237; 3.W.169.238; 3.W.169.239; 3.W.169.154;
3.W.169.157; 3.W.169.166; 3.W.169.169; 3.W.169.172; 3.W.169.175;
3.W.169.240; 3.W.169.244; 3.W.172.228; 3.W.172.229; 3.W.172.230;
20 3.W.172.231; 3.W.172.236; 3.W.172.237; 3.W.172.238; 3.W.172.239;
3.W.172.154; 3.W.172.157; 3.W.172.166; 3.W.172.169; 3.W.172.172;
3.W.172.175; 3.W.172.240; 3.W.172.244; 3.W.175.228; 3.W.175.229;
3.W.175.230; 3.W.175.231; 3.W.175.236; 3.W.175.237; 3.W.175.238;
3.W.175.239; 3.W.175.154; 3.W.175.157; 3.W.175.166; 3.W.175.169;
25 3.W.175.172; 3.W.175.175; 3.W.175.240; 3.W.175.244; 3.W.240.228;
3.W.240.229; 3.W.240.230; 3.W.240.231; 3.W.240.236; 3.W.240.237;
3.W.240.238; 3.W.240.239; 3.W.240.154; 3.W.240.157; 3.W.240.166;
3.W.240.169; 3.W.240.172; 3.W.240.175; 3.W.240.240; 3.W.240.244;
3.W.244.228; 3.W.244.229; 3.W.244.230; 3.W.244.231; 3.W.244.236;
30 3.W.244.237; 3.W.244.238; 3.W.244.239; 3.W.244.154; 3.W.244.157;

3.W.244.166; 3.W.244.169; 3.W.244.172; 3.W.244.175; 3.W.244.240;
3.W.244.244;

Prodrugs of 3.Y

- 5 3.Y.228.228; 3.Y.228.229; 3.Y.228.230; 3.Y.228.231; 3.Y.228.236;
3.Y.228.237; 3.Y.228.238; 3.Y.228.239; 3.Y.228.154; 3.Y.228.157; 3.Y.228.166;
3.Y.228.169; 3.Y.228.172; 3.Y.228.175; 3.Y.228.240; 3.Y.228.244; 3.Y.229.228;
3.Y.229.229; 3.Y.229.230; 3.Y.229.231; 3.Y.229.236; 3.Y.229.237; 3.Y.229.238;
3.Y.229.239; 3.Y.229.154; 3.Y.229.157; 3.Y.229.166; 3.Y.229.169; 3.Y.229.172;
10 3.Y.229.175; 3.Y.229.240; 3.Y.229.244; 3.Y.230.228; 3.Y.230.229; 3.Y.230.230;
3.Y.230.231; 3.Y.230.236; 3.Y.230.237; 3.Y.230.238; 3.Y.230.239; 3.Y.230.154;
3.Y.230.157; 3.Y.230.166; 3.Y.230.169; 3.Y.230.172; 3.Y.230.175; 3.Y.230.240;
3.Y.230.244; 3.Y.231.228; 3.Y.231.229; 3.Y.231.230; 3.Y.231.231; 3.Y.231.236;
3.Y.231.237; 3.Y.231.238; 3.Y.231.239; 3.Y.231.154; 3.Y.231.157; 3.Y.231.166;
15 3.Y.231.169; 3.Y.231.172; 3.Y.231.175; 3.Y.231.240; 3.Y.231.244; 3.Y.236.228;
3.Y.236.229; 3.Y.236.230; 3.Y.236.231; 3.Y.236.236; 3.Y.236.237; 3.Y.236.238;
3.Y.236.239; 3.Y.236.154; 3.Y.236.157; 3.Y.236.166; 3.Y.236.169; 3.Y.236.172;
3.Y.236.175; 3.Y.236.240; 3.Y.236.244; 3.Y.237.228; 3.Y.237.229; 3.Y.237.230;
3.Y.237.231; 3.Y.237.236; 3.Y.237.237; 3.Y.237.238; 3.Y.237.239; 3.Y.237.154;
20 3.Y.237.157; 3.Y.237.166; 3.Y.237.169; 3.Y.237.172; 3.Y.237.175; 3.Y.237.240;
3.Y.237.244; 3.Y.238.228; 3.Y.238.229; 3.Y.238.230; 3.Y.238.231; 3.Y.238.236;
3.Y.238.237; 3.Y.238.238; 3.Y.238.239; 3.Y.238.154; 3.Y.238.157; 3.Y.238.166;
3.Y.238.169; 3.Y.238.172; 3.Y.238.175; 3.Y.238.240; 3.Y.238.244; 3.Y.239.228;
3.Y.239.229; 3.Y.239.230; 3.Y.239.231; 3.Y.239.236; 3.Y.239.237; 3.Y.239.238;
25 3.Y.239.239; 3.Y.239.154; 3.Y.239.157; 3.Y.239.166; 3.Y.239.169; 3.Y.239.172;
3.Y.239.175; 3.Y.239.240; 3.Y.239.244; 3.Y.154.228; 3.Y.154.229; 3.Y.154.230;
3.Y.154.231; 3.Y.154.236; 3.Y.154.237; 3.Y.154.238; 3.Y.154.239; 3.Y.154.154;
3.Y.154.157; 3.Y.154.166; 3.Y.154.169; 3.Y.154.172; 3.Y.154.175; 3.Y.154.240;
3.Y.154.244; 3.Y.157.228; 3.Y.157.229; 3.Y.157.230; 3.Y.157.231; 3.Y.157.236;
30 3.Y.157.237; 3.Y.157.238; 3.Y.157.239; 3.Y.157.154; 3.Y.157.157; 3.Y.157.166;

3.Y.157.169; 3.Y.157.172; 3.Y.157.175; 3.Y.157.240; 3.Y.157.244; 3.Y.166.228;
 3.Y.166.229; 3.Y.166.230; 3.Y.166.231; 3.Y.166.236; 3.Y.166.237; 3.Y.166.238;
 3.Y.166.239; 3.Y.166.154; 3.Y.166.157; 3.Y.166.166; 3.Y.166.169; 3.Y.166.172;
 3.Y.166.175; 3.Y.166.240; 3.Y.166.244; 3.Y.169.228; 3.Y.169.229; 3.Y.169.230;
 5 3.Y.169.231; 3.Y.169.236; 3.Y.169.237; 3.Y.169.238; 3.Y.169.239; 3.Y.169.154;
 3.Y.169.157; 3.Y.169.166; 3.Y.169.169; 3.Y.169.172; 3.Y.169.175; 3.Y.169.240;
 3.Y.169.244; 3.Y.172.228; 3.Y.172.229; 3.Y.172.230; 3.Y.172.231; 3.Y.172.236;
 3.Y.172.237; 3.Y.172.238; 3.Y.172.239; 3.Y.172.154; 3.Y.172.157; 3.Y.172.166;
 3.Y.172.169; 3.Y.172.172; 3.Y.172.175; 3.Y.172.240; 3.Y.172.244; 3.Y.175.228;
 10 3.Y.175.229; 3.Y.175.230; 3.Y.175.231; 3.Y.175.236; 3.Y.175.237; 3.Y.175.238;
 3.Y.175.239; 3.Y.175.154; 3.Y.175.157; 3.Y.175.166; 3.Y.175.169; 3.Y.175.172;
 3.Y.175.175; 3.Y.175.240; 3.Y.175.244; 3.Y.240.228; 3.Y.240.229; 3.Y.240.230;
 3.Y.240.231; 3.Y.240.236; 3.Y.240.237; 3.Y.240.238; 3.Y.240.239; 3.Y.240.154;
 3.Y.240.157; 3.Y.240.166; 3.Y.240.169; 3.Y.240.172; 3.Y.240.175; 3.Y.240.240;
 15 3.Y.240.244; 3.Y.244.228; 3.Y.244.229; 3.Y.244.230; 3.Y.244.231; 3.Y.244.236;
 3.Y.244.237; 3.Y.244.238; 3.Y.244.239; 3.Y.244.154; 3.Y.244.157; 3.Y.244.166;
 3.Y.244.169; 3.Y.244.172; 3.Y.244.175; 3.Y.244.240; 3.Y.244.244;

Prodrugs of 4.B

20 4.B.228.228; 4.B.228.229; 4.B.228.230; 4.B.228.231; 4.B.228.236;
 4.B.228.237; 4.B.228.238; 4.B.228.239; 4.B.228.154; 4.B.228.157; 4.B.228.166;
 4.B.228.169; 4.B.228.172; 4.B.228.175; 4.B.228.240; 4.B.228.244; 4.B.229.228;
 4.B.229.229; 4.B.229.230; 4.B.229.231; 4.B.229.236; 4.B.229.237; 4.B.229.238;
 4.B.229.239; 4.B.229.154; 4.B.229.157; 4.B.229.166; 4.B.229.169; 4.B.229.172;
 25 4.B.229.175; 4.B.229.240; 4.B.229.244; 4.B.230.228; 4.B.230.229; 4.B.230.230;
 4.B.230.231; 4.B.230.236; 4.B.230.237; 4.B.230.238; 4.B.230.239; 4.B.230.154;
 4.B.230.157; 4.B.230.166; 4.B.230.169; 4.B.230.172; 4.B.230.175; 4.B.230.240;
 4.B.230.244; 4.B.231.228; 4.B.231.229; 4.B.231.230; 4.B.231.231; 4.B.231.236;
 4.B.231.237; 4.B.231.238; 4.B.231.239; 4.B.231.154; 4.B.231.157; 4.B.231.166;
 30 4.B.231.169; 4.B.231.172; 4.B.231.175; 4.B.231.240; 4.B.231.244; 4.B.236.228;

4.B.236.229; 4.B.236.230; 4.B.236.231; 4.B.236.236; 4.B.236.237; 4.B.236.238;
4.B.236.239; 4.B.236.154; 4.B.236.157; 4.B.236.166; 4.B.236.169; 4.B.236.172;
4.B.236.175; 4.B.236.240; 4.B.236.244; 4.B.237.228; 4.B.237.229; 4.B.237.230;
4.B.237.231; 4.B.237.236; 4.B.237.237; 4.B.237.238; 4.B.237.239; 4.B.237.154;
5 4.B.237.157; 4.B.237.166; 4.B.237.169; 4.B.237.172; 4.B.237.175; 4.B.237.240;
4.B.237.244; 4.B.238.228; 4.B.238.229; 4.B.238.230; 4.B.238.231; 4.B.238.236;
4.B.238.237; 4.B.238.238; 4.B.238.239; 4.B.238.154; 4.B.238.157; 4.B.238.166;
4.B.238.169; 4.B.238.172; 4.B.238.175; 4.B.238.240; 4.B.238.244; 4.B.239.228;
4.B.239.229; 4.B.239.230; 4.B.239.231; 4.B.239.236; 4.B.239.237; 4.B.239.238;
10 4.B.239.239; 4.B.239.154; 4.B.239.157; 4.B.239.166; 4.B.239.169; 4.B.239.172;
4.B.239.175; 4.B.239.240; 4.B.239.244; 4.B.154.228; 4.B.154.229; 4.B.154.230;
4.B.154.231; 4.B.154.236; 4.B.154.237; 4.B.154.238; 4.B.154.239; 4.B.154.154;
4.B.154.157; 4.B.154.166; 4.B.154.169; 4.B.154.172; 4.B.154.175; 4.B.154.240;
4.B.154.244; 4.B.157.228; 4.B.157.229; 4.B.157.230; 4.B.157.231; 4.B.157.236;
15 4.B.157.237; 4.B.157.238; 4.B.157.239; 4.B.157.154; 4.B.157.157; 4.B.157.166;
4.B.157.169; 4.B.157.172; 4.B.157.175; 4.B.157.240; 4.B.157.244; 4.B.166.228;
4.B.166.229; 4.B.166.230; 4.B.166.231; 4.B.166.236; 4.B.166.237; 4.B.166.238;
4.B.166.239; 4.B.166.154; 4.B.166.157; 4.B.166.166; 4.B.166.169; 4.B.166.172;
4.B.166.175; 4.B.166.240; 4.B.166.244; 4.B.169.228; 4.B.169.229; 4.B.169.230;
20 4.B.169.231; 4.B.169.236; 4.B.169.237; 4.B.169.238; 4.B.169.239; 4.B.169.154;
4.B.169.157; 4.B.169.166; 4.B.169.169; 4.B.169.172; 4.B.169.175; 4.B.169.240;
4.B.169.244; 4.B.172.228; 4.B.172.229; 4.B.172.230; 4.B.172.231; 4.B.172.236;
4.B.172.237; 4.B.172.238; 4.B.172.239; 4.B.172.154; 4.B.172.157; 4.B.172.166;
4.B.172.169; 4.B.172.172; 4.B.172.175; 4.B.172.240; 4.B.172.244; 4.B.175.228;
25 4.B.175.229; 4.B.175.230; 4.B.175.231; 4.B.175.236; 4.B.175.237; 4.B.175.238;
4.B.175.239; 4.B.175.154; 4.B.175.157; 4.B.175.166; 4.B.175.169; 4.B.175.172;
4.B.175.175; 4.B.175.240; 4.B.175.244; 4.B.240.228; 4.B.240.229; 4.B.240.230;
4.B.240.231; 4.B.240.236; 4.B.240.237; 4.B.240.238; 4.B.240.239; 4.B.240.154;
4.B.240.157; 4.B.240.166; 4.B.240.169; 4.B.240.172; 4.B.240.175; 4.B.240.240;
30 4.B.240.244; 4.B.244.228; 4.B.244.229; 4.B.244.230; 4.B.244.231; 4.B.244.236;

4.B.244.237; 4.B.244.238; 4.B.244.239; 4.B.244.154; 4.B.244.157; 4.B.244.166;
4.B.244.169; 4.B.244.172; 4.B.244.175; 4.B.244.240; 4.B.244.244;

Prodrugs of 4.D

- 5 4.D.228.228; 4.D.228.229; 4.D.228.230; 4.D.228.231; 4.D.228.236;
4.D.228.237; 4.D.228.238; 4.D.228.239; 4.D.228.154; 4.D.228.157;
4.D.228.166; 4.D.228.169; 4.D.228.172; 4.D.228.175; 4.D.228.240;
4.D.228.244; 4.D.229.228; 4.D.229.229; 4.D.229.230; 4.D.229.231;
4.D.229.236; 4.D.229.237; 4.D.229.238; 4.D.229.239; 4.D.229.154;
10 4.D.229.157; 4.D.229.166; 4.D.229.169; 4.D.229.172; 4.D.229.175;
4.D.229.240; 4.D.229.244; 4.D.230.228; 4.D.230.229; 4.D.230.230;
4.D.230.231; 4.D.230.236; 4.D.230.237; 4.D.230.238; 4.D.230.239;
4.D.230.154; 4.D.230.157; 4.D.230.166; 4.D.230.169; 4.D.230.172;
4.D.230.175; 4.D.230.240; 4.D.230.244; 4.D.231.228; 4.D.231.229;
15 4.D.231.230; 4.D.231.231; 4.D.231.236; 4.D.231.237; 4.D.231.238;
4.D.231.239; 4.D.231.154; 4.D.231.157; 4.D.231.166; 4.D.231.169;
4.D.231.172; 4.D.231.175; 4.D.231.240; 4.D.231.244; 4.D.236.228;
4.D.236.229; 4.D.236.230; 4.D.236.231; 4.D.236.236; 4.D.236.237;
4.D.236.238; 4.D.236.239; 4.D.236.154; 4.D.236.157; 4.D.236.166;
20 4.D.236.169; 4.D.236.172; 4.D.236.175; 4.D.236.240; 4.D.236.244;
4.D.237.228; 4.D.237.229; 4.D.237.230; 4.D.237.231; 4.D.237.236;
4.D.237.237; 4.D.237.238; 4.D.237.239; 4.D.237.154; 4.D.237.157;
4.D.237.166; 4.D.237.169; 4.D.237.172; 4.D.237.175; 4.D.237.240;
4.D.237.244; 4.D.238.228; 4.D.238.229; 4.D.238.230; 4.D.238.231;
25 4.D.238.236; 4.D.238.237; 4.D.238.238; 4.D.238.239; 4.D.238.154;
4.D.238.157; 4.D.238.166; 4.D.238.169; 4.D.238.172; 4.D.238.175;
4.D.238.240; 4.D.238.244; 4.D.239.228; 4.D.239.229; 4.D.239.230;
4.D.239.231; 4.D.239.236; 4.D.239.237; 4.D.239.238; 4.D.239.239;
4.D.239.154; 4.D.239.157; 4.D.239.166; 4.D.239.169; 4.D.239.172;
30 4.D.239.175; 4.D.239.240; 4.D.239.244; 4.D.154.228; 4.D.154.229;

- 4.D.154.230; 4.D.154.231; 4.D.154.236; 4.D.154.237; 4.D.154.238;
4.D.154.239; 4.D.154.154; 4.D.154.157; 4.D.154.166; 4.D.154.169;
4.D.154.172; 4.D.154.175; 4.D.154.240; 4.D.154.244; 4.D.157.228;
4.D.157.229; 4.D.157.230; 4.D.157.231; 4.D.157.236; 4.D.157.237;
5 4.D.157.238; 4.D.157.239; 4.D.157.154; 4.D.157.157; 4.D.157.166;
4.D.157.169; 4.D.157.172; 4.D.157.175; 4.D.157.240; 4.D.157.244;
4.D.166.228; 4.D.166.229; 4.D.166.230; 4.D.166.231; 4.D.166.236;
4.D.166.237; 4.D.166.238; 4.D.166.239; 4.D.166.154; 4.D.166.157;
4.D.166.166; 4.D.166.169; 4.D.166.172; 4.D.166.175; 4.D.166.240;
10 4.D.166.244; 4.D.169.228; 4.D.169.229; 4.D.169.230; 4.D.169.231;
4.D.169.236; 4.D.169.237; 4.D.169.238; 4.D.169.239; 4.D.169.154;
4.D.169.157; 4.D.169.166; 4.D.169.169; 4.D.169.172; 4.D.169.175;
4.D.169.240; 4.D.169.244; 4.D.172.228; 4.D.172.229; 4.D.172.230;
4.D.172.231; 4.D.172.236; 4.D.172.237; 4.D.172.238; 4.D.172.239;
15 4.D.172.154; 4.D.172.157; 4.D.172.166; 4.D.172.169; 4.D.172.172;
4.D.172.175; 4.D.172.240; 4.D.172.244; 4.D.175.228; 4.D.175.229;
4.D.175.230; 4.D.175.231; 4.D.175.236; 4.D.175.237; 4.D.175.238;
4.D.175.239; 4.D.175.154; 4.D.175.157; 4.D.175.166; 4.D.175.169;
4.D.175.172; 4.D.175.175; 4.D.175.240; 4.D.175.244; 4.D.240.228;
20 4.D.240.229; 4.D.240.230; 4.D.240.231; 4.D.240.236; 4.D.240.237;
4.D.240.238; 4.D.240.239; 4.D.240.154; 4.D.240.157; 4.D.240.166;
4.D.240.169; 4.D.240.172; 4.D.240.175; 4.D.240.240; 4.D.240.244;
4.D.244.228; 4.D.244.229; 4.D.244.230; 4.D.244.231; 4.D.244.236;
4.D.244.237; 4.D.244.238; 4.D.244.239; 4.D.244.154; 4.D.244.157;
25 4.D.244.166; 4.D.244.169; 4.D.244.172; 4.D.244.175; 4.D.244.240;
4.D.244.244;

Prodrugs of 4.E

- 4.E.228.228; 4.E.228.229; 4.E.228.230; 4.E.228.231; 4.E.228.236;
30 4.E.228.237; 4.E.228.238; 4.E.228.239; 4.E.228.154; 4.E.228.157; 4.E.228.166;

4.E.228.169; 4.E.228.172; 4.E.228.175; 4.E.228.240; 4.E.228.244; 4.E.229.228;
4.E.229.229; 4.E.229.230; 4.E.229.231; 4.E.229.236; 4.E.229.237; 4.E.229.238;
4.E.229.239; 4.E.229.154; 4.E.229.157; 4.E.229.166; 4.E.229.169; 4.E.229.172;
4.E.229.175; 4.E.229.240; 4.E.229.244; 4.E.230.228; 4.E.230.229; 4.E.230.230;
5 4.E.230.231; 4.E.230.236; 4.E.230.237; 4.E.230.238; 4.E.230.239; 4.E.230.154;
4.E.230.157; 4.E.230.166; 4.E.230.169; 4.E.230.172; 4.E.230.175; 4.E.230.240;
4.E.230.244; 4.E.231.228; 4.E.231.229; 4.E.231.230; 4.E.231.231; 4.E.231.236;
4.E.231.237; 4.E.231.238; 4.E.231.239; 4.E.231.154; 4.E.231.157; 4.E.231.166;
4.E.231.169; 4.E.231.172; 4.E.231.175; 4.E.231.240; 4.E.231.244; 4.E.236.228;
10 4.E.236.229; 4.E.236.230; 4.E.236.231; 4.E.236.236; 4.E.236.237; 4.E.236.238;
4.E.236.239; 4.E.236.154; 4.E.236.157; 4.E.236.166; 4.E.236.169; 4.E.236.172;
4.E.236.175; 4.E.236.240; 4.E.236.244; 4.E.237.228; 4.E.237.229; 4.E.237.230;
4.E.237.231; 4.E.237.236; 4.E.237.237; 4.E.237.238; 4.E.237.239; 4.E.237.154;
4.E.237.157; 4.E.237.166; 4.E.237.169; 4.E.237.172; 4.E.237.175; 4.E.237.240;
15 4.E.237.244; 4.E.238.228; 4.E.238.229; 4.E.238.230; 4.E.238.231; 4.E.238.236;
4.E.238.237; 4.E.238.238; 4.E.238.239; 4.E.238.154; 4.E.238.157; 4.E.238.166;
4.E.238.169; 4.E.238.172; 4.E.238.175; 4.E.238.240; 4.E.238.244; 4.E.239.228;
4.E.239.229; 4.E.239.230; 4.E.239.231; 4.E.239.236; 4.E.239.237; 4.E.239.238;
4.E.239.239; 4.E.239.154; 4.E.239.157; 4.E.239.166; 4.E.239.169; 4.E.239.172;
20 4.E.239.175; 4.E.239.240; 4.E.239.244; 4.E.154.228; 4.E.154.229; 4.E.154.230;
4.E.154.231; 4.E.154.236; 4.E.154.237; 4.E.154.238; 4.E.154.239; 4.E.154.154;
4.E.154.157; 4.E.154.166; 4.E.154.169; 4.E.154.172; 4.E.154.175; 4.E.154.240;
4.E.154.244; 4.E.157.228; 4.E.157.229; 4.E.157.230; 4.E.157.231; 4.E.157.236;
4.E.157.237; 4.E.157.238; 4.E.157.239; 4.E.157.154; 4.E.157.157; 4.E.157.166;
25 4.E.157.169; 4.E.157.172; 4.E.157.175; 4.E.157.240; 4.E.157.244; 4.E.166.228;
4.E.166.229; 4.E.166.230; 4.E.166.231; 4.E.166.236; 4.E.166.237; 4.E.166.238;
4.E.166.239; 4.E.166.154; 4.E.166.157; 4.E.166.166; 4.E.166.169; 4.E.166.172;
4.E.166.175; 4.E.166.240; 4.E.166.244; 4.E.169.228; 4.E.169.229; 4.E.169.230;
4.E.169.231; 4.E.169.236; 4.E.169.237; 4.E.169.238; 4.E.169.239; 4.E.169.154;
30 4.E.169.157; 4.E.169.166; 4.E.169.169; 4.E.169.172; 4.E.169.175; 4.E.169.240;

4.E.169.244; 4.E.172.228; 4.E.172.229; 4.E.172.230; 4.E.172.231; 4.E.172.236;
4.E.172.237; 4.E.172.238; 4.E.172.239; 4.E.172.154; 4.E.172.157; 4.E.172.166;
4.E.172.169; 4.E.172.172; 4.E.172.175; 4.E.172.240; 4.E.172.244; 4.E.175.228;
4.E.175.229; 4.E.175.230; 4.E.175.231; 4.E.175.236; 4.E.175.237; 4.E.175.238;
5 4.E.175.239; 4.E.175.154; 4.E.175.157; 4.E.175.166; 4.E.175.169; 4.E.175.172;
4.E.175.175; 4.E.175.240; 4.E.175.244; 4.E.240.228; 4.E.240.229; 4.E.240.230;
4.E.240.231; 4.E.240.236; 4.E.240.237; 4.E.240.238; 4.E.240.239; 4.E.240.154;
4.E.240.157; 4.E.240.166; 4.E.240.169; 4.E.240.172; 4.E.240.175; 4.E.240.240;
4.E.240.244; 4.E.244.228; 4.E.244.229; 4.E.244.230; 4.E.244.231; 4.E.244.236;
10 4.E.244.237; 4.E.244.238; 4.E.244.239; 4.E.244.154; 4.E.244.157; 4.E.244.166;
4.E.244.169; 4.E.244.172; 4.E.244.175; 4.E.244.240; 4.E.244.244;

Prodrugs of 4.G

4.G.228.228; 4.G.228.229; 4.G.228.230; 4.G.228.231; 4.G.228.236;
15 4.G.228.237; 4.G.228.238; 4.G.228.239; 4.G.228.154; 4.G.228.157;
4.G.228.166; 4.G.228.169; 4.G.228.172; 4.G.228.175; 4.G.228.240;
4.G.228.244; 4.G.229.228; 4.G.229.229; 4.G.229.230; 4.G.229.231;
4.G.229.236; 4.G.229.237; 4.G.229.238; 4.G.229.239; 4.G.229.154;
4.G.229.157; 4.G.229.166; 4.G.229.169; 4.G.229.172; 4.G.229.175;
20 4.G.229.240; 4.G.229.244; 4.G.230.228; 4.G.230.229; 4.G.230.230;
4.G.230.231; 4.G.230.236; 4.G.230.237; 4.G.230.238; 4.G.230.239;
4.G.230.154; 4.G.230.157; 4.G.230.166; 4.G.230.169; 4.G.230.172;
4.G.230.175; 4.G.230.240; 4.G.230.244; 4.G.231.228; 4.G.231.229;
4.G.231.230; 4.G.231.231; 4.G.231.236; 4.G.231.237; 4.G.231.238;
25 4.G.231.239; 4.G.231.154; 4.G.231.157; 4.G.231.166; 4.G.231.169;
4.G.231.172; 4.G.231.175; 4.G.231.240; 4.G.231.244; 4.G.236.228;
4.G.236.229; 4.G.236.230; 4.G.236.231; 4.G.236.236; 4.G.236.237;
4.G.236.238; 4.G.236.239; 4.G.236.154; 4.G.236.157; 4.G.236.166;
4.G.236.169; 4.G.236.172; 4.G.236.175; 4.G.236.240; 4.G.236.244;
30 4.G.237.228; 4.G.237.229; 4.G.237.230; 4.G.237.231; 4.G.237.236;

- 4.G.237.237; 4.G.237.238; 4.G.237.239; 4.G.237.154; 4.G.237.157;
4.G.237.166; 4.G.237.169; 4.G.237.172; 4.G.237.175; 4.G.237.240;
4.G.237.244; 4.G.238.228; 4.G.238.229; 4.G.238.230; 4.G.238.231;
4.G.238.236; 4.G.238.237; 4.G.238.238; 4.G.238.239; 4.G.238.154;
5 4.G.238.157; 4.G.238.166; 4.G.238.169; 4.G.238.172; 4.G.238.175;
4.G.238.240; 4.G.238.244; 4.G.239.228; 4.G.239.229; 4.G.239.230;
4.G.239.231; 4.G.239.236; 4.G.239.237; 4.G.239.238; 4.G.239.239;
4.G.239.154; 4.G.239.157; 4.G.239.166; 4.G.239.169; 4.G.239.172;
4.G.239.175; 4.G.239.240; 4.G.239.244; 4.G.154.228; 4.G.154.229;
10 4.G.154.230; 4.G.154.231; 4.G.154.236; 4.G.154.237; 4.G.154.238;
4.G.154.239; 4.G.154.154; 4.G.154.157; 4.G.154.166; 4.G.154.169;
4.G.154.172; 4.G.154.175; 4.G.154.240; 4.G.154.244; 4.G.157.228;
4.G.157.229; 4.G.157.230; 4.G.157.231; 4.G.157.236; 4.G.157.237;
4.G.157.238; 4.G.157.239; 4.G.157.154; 4.G.157.157; 4.G.157.166;
15 4.G.157.169; 4.G.157.172; 4.G.157.175; 4.G.157.240; 4.G.157.244;
4.G.166.228; 4.G.166.229; 4.G.166.230; 4.G.166.231; 4.G.166.236;
4.G.166.237; 4.G.166.238; 4.G.166.239; 4.G.166.154; 4.G.166.157;
4.G.166.166; 4.G.166.169; 4.G.166.172; 4.G.166.175; 4.G.166.240;
4.G.166.244; 4.G.169.228; 4.G.169.229; 4.G.169.230; 4.G.169.231;
20 4.G.169.236; 4.G.169.237; 4.G.169.238; 4.G.169.239; 4.G.169.154;
4.G.169.157; 4.G.169.166; 4.G.169.169; 4.G.169.172; 4.G.169.175;
4.G.169.240; 4.G.169.244; 4.G.172.228; 4.G.172.229; 4.G.172.230;
4.G.172.231; 4.G.172.236; 4.G.172.237; 4.G.172.238; 4.G.172.239;
4.G.172.154; 4.G.172.157; 4.G.172.166; 4.G.172.169; 4.G.172.172;
25 4.G.172.175; 4.G.172.240; 4.G.172.244; 4.G.175.228; 4.G.175.229;
4.G.175.230; 4.G.175.231; 4.G.175.236; 4.G.175.237; 4.G.175.238;
4.G.175.239; 4.G.175.154; 4.G.175.157; 4.G.175.166; 4.G.175.169;
4.G.175.172; 4.G.175.175; 4.G.175.240; 4.G.175.244; 4.G.240.228;
4.G.240.229; 4.G.240.230; 4.G.240.231; 4.G.240.236; 4.G.240.237;
30 4.G.240.238; 4.G.240.239; 4.G.240.154; 4.G.240.157; 4.G.240.166;

4.G.240.169; 4.G.240.172; 4.G.240.175; 4.G.240.240; 4.G.240.244;
 4.G.244.228; 4.G.244.229; 4.G.244.230; 4.G.244.231; 4.G.244.236;
 4.G.244.237; 4.G.244.238; 4.G.244.239; 4.G.244.154; 4.G.244.157;
 4.G.244.166; 4.G.244.169; 4.G.244.172; 4.G.244.175; 4.G.244.240;
 5 4.G.244.244;

Prodrugs of 4.I

4.I.228.228; 4.I.228.229; 4.I.228.230; 4.I.228.231; 4.I.228.236; 4.I.228.237;
 4.I.228.238; 4.I.228.239; 4.I.228.154; 4.I.228.157; 4.I.228.166; 4.I.228.169;
 10 4.I.228.172; 4.I.228.175; 4.I.228.240; 4.I.228.244; 4.I.229.228; 4.I.229.229;
 4.I.229.230; 4.I.229.231; 4.I.229.236; 4.I.229.237; 4.I.229.238; 4.I.229.239;
 4.I.229.154; 4.I.229.157; 4.I.229.166; 4.I.229.169; 4.I.229.172; 4.I.229.175;
 4.I.229.240; 4.I.229.244; 4.I.230.228; 4.I.230.229; 4.I.230.230; 4.I.230.231;
 4.I.230.236; 4.I.230.237; 4.I.230.238; 4.I.230.239; 4.I.230.154; 4.I.230.157;
 15 4.I.230.166; 4.I.230.169; 4.I.230.172; 4.I.230.175; 4.I.230.240; 4.I.230.244;
 4.I.231.228; 4.I.231.229; 4.I.231.230; 4.I.231.231; 4.I.231.236; 4.I.231.237;
 4.I.231.238; 4.I.231.239; 4.I.231.154; 4.I.231.157; 4.I.231.166; 4.I.231.169;
 4.I.231.172; 4.I.231.175; 4.I.231.240; 4.I.231.244; 4.I.236.228; 4.I.236.229;
 4.I.236.230; 4.I.236.231; 4.I.236.236; 4.I.236.237; 4.I.236.238; 4.I.236.239;
 20 4.I.236.154; 4.I.236.157; 4.I.236.166; 4.I.236.169; 4.I.236.172; 4.I.236.175;
 4.I.236.240; 4.I.236.244; 4.I.237.228; 4.I.237.229; 4.I.237.230; 4.I.237.231;
 4.I.237.236; 4.I.237.237; 4.I.237.238; 4.I.237.239; 4.I.237.154; 4.I.237.157;
 4.I.237.166; 4.I.237.169; 4.I.237.172; 4.I.237.175; 4.I.237.240; 4.I.237.244;
 4.I.238.228; 4.I.238.229; 4.I.238.230; 4.I.238.231; 4.I.238.236; 4.I.238.237;
 25 4.I.238.238; 4.I.238.239; 4.I.238.154; 4.I.238.157; 4.I.238.166; 4.I.238.169;
 4.I.238.172; 4.I.238.175; 4.I.238.240; 4.I.238.244; 4.I.239.228; 4.I.239.229;
 4.I.239.230; 4.I.239.231; 4.I.239.236; 4.I.239.237; 4.I.239.238; 4.I.239.239;
 4.I.239.154; 4.I.239.157; 4.I.239.166; 4.I.239.169; 4.I.239.172; 4.I.239.175;
 4.I.239.240; 4.I.239.244; 4.I.154.228; 4.I.154.229; 4.I.154.230; 4.I.154.231;
 30 4.I.154.236; 4.I.154.237; 4.I.154.238; 4.I.154.239; 4.I.154.154; 4.I.154.157;

- 4.I.154.166; 4.I.154.169; 4.I.154.172; 4.I.154.175; 4.I.154.240; 4.I.154.244;
4.I.157.228; 4.I.157.229; 4.I.157.230; 4.I.157.231; 4.I.157.236; 4.I.157.237;
4.I.157.238; 4.I.157.239; 4.I.157.154; 4.I.157.157; 4.I.157.166; 4.I.157.169;
4.I.157.172; 4.I.157.175; 4.I.157.240; 4.I.157.244; 4.I.166.228; 4.I.166.229;
5 4.I.166.230; 4.I.166.231; 4.I.166.236; 4.I.166.237; 4.I.166.238; 4.I.166.239;
4.I.166.154; 4.I.166.157; 4.I.166.166; 4.I.166.169; 4.I.166.172; 4.I.166.175;
4.I.166.240; 4.I.166.244; 4.I.169.228; 4.I.169.229; 4.I.169.230; 4.I.169.231;
4.I.169.236; 4.I.169.237; 4.I.169.238; 4.I.169.239; 4.I.169.154; 4.I.169.157;
4.I.169.166; 4.I.169.169; 4.I.169.172; 4.I.169.175; 4.I.169.240; 4.I.169.244;
10 4.I.172.228; 4.I.172.229; 4.I.172.230; 4.I.172.231; 4.I.172.236; 4.I.172.237;
4.I.172.238; 4.I.172.239; 4.I.172.154; 4.I.172.157; 4.I.172.166; 4.I.172.169;
4.I.172.172; 4.I.172.175; 4.I.172.240; 4.I.172.244; 4.I.175.228; 4.I.175.229;
4.I.175.230; 4.I.175.231; 4.I.175.236; 4.I.175.237; 4.I.175.238; 4.I.175.239;
4.I.175.154; 4.I.175.157; 4.I.175.166; 4.I.175.169; 4.I.175.172; 4.I.175.175;
15 4.I.175.240; 4.I.175.244; 4.I.240.228; 4.I.240.229; 4.I.240.230; 4.I.240.231;
4.I.240.236; 4.I.240.237; 4.I.240.238; 4.I.240.239; 4.I.240.154; 4.I.240.157;
4.I.240.166; 4.I.240.169; 4.I.240.172; 4.I.240.175; 4.I.240.240; 4.I.240.244;
4.I.244.228; 4.I.244.229; 4.I.244.230; 4.I.244.231; 4.I.244.236; 4.I.244.237;
4.I.244.238; 4.I.244.239; 4.I.244.154; 4.I.244.157; 4.I.244.166; 4.I.244.169;
20 4.I.244.172; 4.I.244.175; 4.I.244.240; 4.I.244.244;

Prodrugs of 4.I

- 4.J.228.228; 4.J.228.229; 4.J.228.230; 4.J.228.231; 4.J.228.236; 4.J.228.237;
4.J.228.238; 4.J.228.239; 4.J.228.154; 4.J.228.157; 4.J.228.166; 4.J.228.169;
25 4.J.228.172; 4.J.228.175; 4.J.228.240; 4.J.228.244; 4.J.229.228; 4.J.229.229;
4.J.229.230; 4.J.229.231; 4.J.229.236; 4.J.229.237; 4.J.229.238; 4.J.229.239;
4.J.229.154; 4.J.229.157; 4.J.229.166; 4.J.229.169; 4.J.229.172; 4.J.229.175;
4.J.229.240; 4.J.229.244; 4.J.230.228; 4.J.230.229; 4.J.230.230; 4.J.230.231;
4.J.230.236; 4.J.230.237; 4.J.230.238; 4.J.230.239; 4.J.230.154; 4.J.230.157;
30 4.J.230.166; 4.J.230.169; 4.J.230.172; 4.J.230.175; 4.J.230.240; 4.J.230.244;

4.J.231.228; 4.J.231.229; 4.J.231.230; 4.J.231.231; 4.J.231.236; 4.J.231.237;
 4.J.231.238; 4.J.231.239; 4.J.231.154; 4.J.231.157; 4.J.231.166; 4.J.231.169;
 4.J.231.172; 4.J.231.175; 4.J.231.240; 4.J.231.244; 4.J.236.228; 4.J.236.229;
 4.J.236.230; 4.J.236.231; 4.J.236.236; 4.J.236.237; 4.J.236.238; 4.J.236.239;
 5 4.J.236.154; 4.J.236.157; 4.J.236.166; 4.J.236.169; 4.J.236.172; 4.J.236.175;
 4.J.236.240; 4.J.236.244; 4.J.237.228; 4.J.237.229; 4.J.237.230; 4.J.237.231;
 4.J.237.236; 4.J.237.237; 4.J.237.238; 4.J.237.239; 4.J.237.154; 4.J.237.157;
 4.J.237.166; 4.J.237.169; 4.J.237.172; 4.J.237.175; 4.J.237.240; 4.J.237.244;
 4.J.238.228; 4.J.238.229; 4.J.238.230; 4.J.238.231; 4.J.238.236; 4.J.238.237;
 10 4.J.238.238; 4.J.238.239; 4.J.238.154; 4.J.238.157; 4.J.238.166; 4.J.238.169;
 4.J.238.172; 4.J.238.175; 4.J.238.240; 4.J.238.244; 4.J.239.228; 4.J.239.229;
 4.J.239.230; 4.J.239.231; 4.J.239.236; 4.J.239.237; 4.J.239.238; 4.J.239.239;
 4.J.239.154; 4.J.239.157; 4.J.239.166; 4.J.239.169; 4.J.239.172; 4.J.239.175;
 4.J.239.240; 4.J.239.244; 4.J.154.228; 4.J.154.229; 4.J.154.230; 4.J.154.231;
 15 4.J.154.236; 4.J.154.237; 4.J.154.238; 4.J.154.239; 4.J.154.154; 4.J.154.157;
 4.J.154.166; 4.J.154.169; 4.J.154.172; 4.J.154.175; 4.J.154.240; 4.J.154.244;
 4.J.157.228; 4.J.157.229; 4.J.157.230; 4.J.157.231; 4.J.157.236; 4.J.157.237;
 4.J.157.238; 4.J.157.239; 4.J.157.154; 4.J.157.157; 4.J.157.166; 4.J.157.169;
 4.J.157.172; 4.J.157.175; 4.J.157.240; 4.J.157.244; 4.J.166.228; 4.J.166.229;
 20 4.J.166.230; 4.J.166.231; 4.J.166.236; 4.J.166.237; 4.J.166.238; 4.J.166.239;
 4.J.166.154; 4.J.166.157; 4.J.166.166; 4.J.166.169; 4.J.166.172; 4.J.166.175;
 4.J.166.240; 4.J.166.244; 4.J.169.228; 4.J.169.229; 4.J.169.230; 4.J.169.231;
 4.J.169.236; 4.J.169.237; 4.J.169.238; 4.J.169.239; 4.J.169.154; 4.J.169.157;
 4.J.169.166; 4.J.169.169; 4.J.169.172; 4.J.169.175; 4.J.169.240; 4.J.169.244;
 25 4.J.172.228; 4.J.172.229; 4.J.172.230; 4.J.172.231; 4.J.172.236; 4.J.172.237;
 4.J.172.238; 4.J.172.239; 4.J.172.154; 4.J.172.157; 4.J.172.166; 4.J.172.169;
 4.J.172.172; 4.J.172.175; 4.J.172.240; 4.J.172.244; 4.J.175.228; 4.J.175.229;
 4.J.175.230; 4.J.175.231; 4.J.175.236; 4.J.175.237; 4.J.175.238; 4.J.175.239;
 4.J.175.154; 4.J.175.157; 4.J.175.166; 4.J.175.169; 4.J.175.172; 4.J.175.175;
 30 4.J.175.240; 4.J.175.244; 4.J.240.228; 4.J.240.229; 4.J.240.230; 4.J.240.231;

4.J.240.236; 4.J.240.237; 4.J.240.238; 4.J.240.239; 4.J.240.154; 4.J.240.157;
4.J.240.166; 4.J.240.169; 4.J.240.172; 4.J.240.175; 4.J.240.240; 4.J.240.244;
4.J.244.228; 4.J.244.229; 4.J.244.230; 4.J.244.231; 4.J.244.236; 4.J.244.237;
4.J.244.238; 4.J.244.239; 4.J.244.154; 4.J.244.157; 4.J.244.166; 4.J.244.169;
5 4.J.244.172; 4.J.244.175; 4.J.244.240; 4.J.244.244;

Prodrugs of 4.L

4.L.228.228; 4.L.228.229; 4.L.228.230; 4.L.228.231; 4.L.228.236;
4.L.228.237; 4.L.228.238; 4.L.228.239; 4.L.228.154; 4.L.228.157; 4.L.228.166;
10 4.L.228.169; 4.L.228.172; 4.L.228.175; 4.L.228.240; 4.L.228.244; 4.L.229.228;
4.L.229.229; 4.L.229.230; 4.L.229.231; 4.L.229.236; 4.L.229.237; 4.L.229.238;
4.L.229.239; 4.L.229.154; 4.L.229.157; 4.L.229.166; 4.L.229.169; 4.L.229.172;
4.L.229.175; 4.L.229.240; 4.L.229.244; 4.L.230.228; 4.L.230.229; 4.L.230.230;
4.L.230.231; 4.L.230.236; 4.L.230.237; 4.L.230.238; 4.L.230.239; 4.L.230.154;
15 4.L.230.157; 4.L.230.166; 4.L.230.169; 4.L.230.172; 4.L.230.175; 4.L.230.240;
4.L.230.244; 4.L.231.228; 4.L.231.229; 4.L.231.230; 4.L.231.231; 4.L.231.236;
4.L.231.237; 4.L.231.238; 4.L.231.239; 4.L.231.154; 4.L.231.157; 4.L.231.166;
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20 4.L.236.239; 4.L.236.154; 4.L.236.157; 4.L.236.166; 4.L.236.169; 4.L.236.172;
4.L.236.175; 4.L.236.240; 4.L.236.244; 4.L.237.228; 4.L.237.229; 4.L.237.230;
4.L.237.231; 4.L.237.236; 4.L.237.237; 4.L.237.238; 4.L.237.239; 4.L.237.154;
4.L.237.157; 4.L.237.166; 4.L.237.169; 4.L.237.172; 4.L.237.175; 4.L.237.240;
4.L.237.244; 4.L.238.228; 4.L.238.229; 4.L.238.230; 4.L.238.231; 4.L.238.236;
25 4.L.238.237; 4.L.238.238; 4.L.238.239; 4.L.238.154; 4.L.238.157; 4.L.238.166;
4.L.238.169; 4.L.238.172; 4.L.238.175; 4.L.238.240; 4.L.238.244; 4.L.239.228;
4.L.239.229; 4.L.239.230; 4.L.239.231; 4.L.239.236; 4.L.239.237; 4.L.239.238;
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4.L.239.175; 4.L.239.240; 4.L.239.244; 4.L.154.228; 4.L.154.229; 4.L.154.230;
30 4.L.154.231; 4.L.154.236; 4.L.154.237; 4.L.154.238; 4.L.154.239; 4.L.154.154;

4.L.154.157; 4.L.154.166; 4.L.154.169; 4.L.154.172; 4.L.154.175; 4.L.154.240;
4.L.154.244; 4.L.157.228; 4.L.157.229; 4.L.157.230; 4.L.157.231; 4.L.157.236;
4.L.157.237; 4.L.157.238; 4.L.157.239; 4.L.157.154; 4.L.157.157; 4.L.157.166;
4.L.157.169; 4.L.157.172; 4.L.157.175; 4.L.157.240; 4.L.157.244; 4.L.166.228;
5 4.L.166.229; 4.L.166.230; 4.L.166.231; 4.L.166.236; 4.L.166.237; 4.L.166.238;
4.L.166.239; 4.L.166.154; 4.L.166.157; 4.L.166.166; 4.L.166.169; 4.L.166.172;
4.L.166.175; 4.L.166.240; 4.L.166.244; 4.L.169.228; 4.L.169.229; 4.L.169.230;
4.L.169.231; 4.L.169.236; 4.L.169.237; 4.L.169.238; 4.L.169.239; 4.L.169.154;
4.L.169.157; 4.L.169.166; 4.L.169.169; 4.L.169.172; 4.L.169.175; 4.L.169.240;
10 4.L.169.244; 4.L.172.228; 4.L.172.229; 4.L.172.230; 4.L.172.231; 4.L.172.236;
4.L.172.237; 4.L.172.238; 4.L.172.239; 4.L.172.154; 4.L.172.157; 4.L.172.166;
4.L.172.169; 4.L.172.172; 4.L.172.175; 4.L.172.240; 4.L.172.244; 4.L.175.228;
4.L.175.229; 4.L.175.230; 4.L.175.231; 4.L.175.236; 4.L.175.237; 4.L.175.238;
4.L.175.239; 4.L.175.154; 4.L.175.157; 4.L.175.166; 4.L.175.169; 4.L.175.172;
15 4.L.175.175; 4.L.175.240; 4.L.175.244; 4.L.240.228; 4.L.240.229; 4.L.240.230;
4.L.240.231; 4.L.240.236; 4.L.240.237; 4.L.240.238; 4.L.240.239; 4.L.240.154;
4.L.240.157; 4.L.240.166; 4.L.240.169; 4.L.240.172; 4.L.240.175; 4.L.240.240;
4.L.240.244; 4.L.244.228; 4.L.244.229; 4.L.244.230; 4.L.244.231; 4.L.244.236;
4.L.244.237; 4.L.244.238; 4.L.244.239; 4.L.244.154; 4.L.244.157; 4.L.244.166;
20 4.L.244.169; 4.L.244.172; 4.L.244.175; 4.L.244.240; 4.L.244.244;

Prodrugs of 4.O

4.O.228.228; 4.O.228.229; 4.O.228.230; 4.O.228.231; 4.O.228.236;
4.O.228.237; 4.O.228.238; 4.O.228.239; 4.O.228.154; 4.O.228.157;
25 4.O.228.166; 4.O.228.169; 4.O.228.172; 4.O.228.175; 4.O.228.240;
4.O.228.244; 4.O.229.228; 4.O.229.229; 4.O.229.230; 4.O.229.231;
4.O.229.236; 4.O.229.237; 4.O.229.238; 4.O.229.239; 4.O.229.154;
4.O.229.157; 4.O.229.166; 4.O.229.169; 4.O.229.172; 4.O.229.175;
4.O.229.240; 4.O.229.244; 4.O.230.228; 4.O.230.229; 4.O.230.230;
30 4.O.230.231; 4.O.230.236; 4.O.230.237; 4.O.230.238; 4.O.230.239;

4.O.230.154; 4.O.230.157; 4.O.230.166; 4.O.230.169; 4.O.230.172;
4.O.230.175; 4.O.230.240; 4.O.230.244; 4.O.231.228; 4.O.231.229;
4.O.231.230; 4.O.231.231; 4.O.231.236; 4.O.231.237; 4.O.231.238;
4.O.231.239; 4.O.231.154; 4.O.231.157; 4.O.231.166; 4.O.231.169;
5 4.O.231.172; 4.O.231.175; 4.O.231.240; 4.O.231.244; 4.O.236.228;
4.O.236.229; 4.O.236.230; 4.O.236.231; 4.O.236.236; 4.O.236.237;
4.O.236.238; 4.O.236.239; 4.O.236.154; 4.O.236.157; 4.O.236.166;
4.O.236.169; 4.O.236.172; 4.O.236.175; 4.O.236.240; 4.O.236.244;
4.O.237.228; 4.O.237.229; 4.O.237.230; 4.O.237.231; 4.O.237.236;
10 4.O.237.237; 4.O.237.238; 4.O.237.239; 4.O.237.154; 4.O.237.157;
4.O.237.166; 4.O.237.169; 4.O.237.172; 4.O.237.175; 4.O.237.240;
4.O.237.244; 4.O.238.228; 4.O.238.229; 4.O.238.230; 4.O.238.231;
4.O.238.236; 4.O.238.237; 4.O.238.238; 4.O.238.239; 4.O.238.154;
4.O.238.157; 4.O.238.166; 4.O.238.169; 4.O.238.172; 4.O.238.175;
15 4.O.238.240; 4.O.238.244; 4.O.239.228; 4.O.239.229; 4.O.239.230;
4.O.239.231; 4.O.239.236; 4.O.239.237; 4.O.239.238; 4.O.239.239;
4.O.239.154; 4.O.239.157; 4.O.239.166; 4.O.239.169; 4.O.239.172;
4.O.239.175; 4.O.239.240; 4.O.239.244; 4.O.154.228; 4.O.154.229;
4.O.154.230; 4.O.154.231; 4.O.154.236; 4.O.154.237; 4.O.154.238;
20 4.O.154.239; 4.O.154.154; 4.O.154.157; 4.O.154.166; 4.O.154.169;
4.O.154.172; 4.O.154.175; 4.O.154.240; 4.O.154.244; 4.O.157.228;
4.O.157.229; 4.O.157.230; 4.O.157.231; 4.O.157.236; 4.O.157.237;
4.O.157.238; 4.O.157.239; 4.O.157.154; 4.O.157.157; 4.O.157.166;
4.O.157.169; 4.O.157.172; 4.O.157.175; 4.O.157.240; 4.O.157.244;
25 4.O.166.228; 4.O.166.229; 4.O.166.230; 4.O.166.231; 4.O.166.236;
4.O.166.237; 4.O.166.238; 4.O.166.239; 4.O.166.154; 4.O.166.157;
4.O.166.166; 4.O.166.169; 4.O.166.172; 4.O.166.175; 4.O.166.240;
4.O.166.244; 4.O.169.228; 4.O.169.229; 4.O.169.230; 4.O.169.231;
4.O.169.236; 4.O.169.237; 4.O.169.238; 4.O.169.239; 4.O.169.154;
30 4.O.169.157; 4.O.169.166; 4.O.169.169; 4.O.169.172; 4.O.169.175;

4.O.169.240; 4.O.169.244; 4.O.172.228; 4.O.172.229; 4.O.172.230;
4.O.172.231; 4.O.172.236; 4.O.172.237; 4.O.172.238; 4.O.172.239;
4.O.172.154; 4.O.172.157; 4.O.172.166; 4.O.172.169; 4.O.172.172;
4.O.172.175; 4.O.172.240; 4.O.172.244; 4.O.175.228; 4.O.175.229;
5 4.O.175.230; 4.O.175.231; 4.O.175.236; 4.O.175.237; 4.O.175.238;
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4.O.175.172; 4.O.175.175; 4.O.175.240; 4.O.175.244; 4.O.240.228;
4.O.240.229; 4.O.240.230; 4.O.240.231; 4.O.240.236; 4.O.240.237;
4.O.240.238; 4.O.240.239; 4.O.240.154; 4.O.240.157; 4.O.240.166;
10 4.O.240.169; 4.O.240.172; 4.O.240.175; 4.O.240.240; 4.O.240.244;
4.O.244.228; 4.O.244.229; 4.O.244.230; 4.O.244.231; 4.O.244.236;
4.O.244.237; 4.O.244.238; 4.O.244.239; 4.O.244.154; 4.O.244.157;
4.O.244.166; 4.O.244.169; 4.O.244.172; 4.O.244.175; 4.O.244.240;
4.O.244.244;

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Prodrugs of 4.P

4.P.228.228; 4.P.228.229; 4.P.228.230; 4.P.228.231; 4.P.228.236;
4.P.228.237; 4.P.228.238; 4.P.228.239; 4.P.228.154; 4.P.228.157; 4.P.228.166;
4.P.228.169; 4.P.228.172; 4.P.228.175; 4.P.228.240; 4.P.228.244; 4.P.229.228;
20 4.P.229.229; 4.P.229.230; 4.P.229.231; 4.P.229.236; 4.P.229.237; 4.P.229.238;
4.P.229.239; 4.P.229.154; 4.P.229.157; 4.P.229.166; 4.P.229.169; 4.P.229.172;
4.P.229.175; 4.P.229.240; 4.P.229.244; 4.P.230.228; 4.P.230.229; 4.P.230.230;
4.P.230.231; 4.P.230.236; 4.P.230.237; 4.P.230.238; 4.P.230.239; 4.P.230.154;
4.P.230.157; 4.P.230.166; 4.P.230.169; 4.P.230.172; 4.P.230.175; 4.P.230.240;
25 4.P.230.244; 4.P.231.228; 4.P.231.229; 4.P.231.230; 4.P.231.231; 4.P.231.236;
4.P.231.237; 4.P.231.238; 4.P.231.239; 4.P.231.154; 4.P.231.157; 4.P.231.166;
4.P.231.169; 4.P.231.172; 4.P.231.175; 4.P.231.240; 4.P.231.244; 4.P.236.228;
4.P.236.229; 4.P.236.230; 4.P.236.231; 4.P.236.236; 4.P.236.237; 4.P.236.238;
4.P.236.239; 4.P.236.154; 4.P.236.157; 4.P.236.166; 4.P.236.169; 4.P.236.172;
30 4.P.236.175; 4.P.236.240; 4.P.236.244; 4.P.237.228; 4.P.237.229; 4.P.237.230;

4.P.237.231; 4.P.237.236; 4.P.237.237; 4.P.237.238; 4.P.237.239; 4.P.237.154;
4.P.237.157; 4.P.237.166; 4.P.237.169; 4.P.237.172; 4.P.237.175; 4.P.237.240;
4.P.237.244; 4.P.238.228; 4.P.238.229; 4.P.238.230; 4.P.238.231; 4.P.238.236;
4.P.238.237; 4.P.238.238; 4.P.238.239; 4.P.238.154; 4.P.238.157; 4.P.238.166;
5 4.P.238.169; 4.P.238.172; 4.P.238.175; 4.P.238.240; 4.P.238.244; 4.P.239.228;
4.P.239.229; 4.P.239.230; 4.P.239.231; 4.P.239.236; 4.P.239.237; 4.P.239.238;
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4.P.239.175; 4.P.239.240; 4.P.239.244; 4.P.154.228; 4.P.154.229; 4.P.154.230;
4.P.154.231; 4.P.154.236; 4.P.154.237; 4.P.154.238; 4.P.154.239; 4.P.154.154;
10 4.P.154.157; 4.P.154.166; 4.P.154.169; 4.P.154.172; 4.P.154.175; 4.P.154.240;
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4.P.157.237; 4.P.157.238; 4.P.157.239; 4.P.157.154; 4.P.157.157; 4.P.157.166;
4.P.157.169; 4.P.157.172; 4.P.157.175; 4.P.157.240; 4.P.157.244; 4.P.166.228;
4.P.166.229; 4.P.166.230; 4.P.166.231; 4.P.166.236; 4.P.166.237; 4.P.166.238;
15 4.P.166.239; 4.P.166.154; 4.P.166.157; 4.P.166.166; 4.P.166.169; 4.P.166.172;
4.P.166.175; 4.P.166.240; 4.P.166.244; 4.P.169.228; 4.P.169.229; 4.P.169.230;
4.P.169.231; 4.P.169.236; 4.P.169.237; 4.P.169.238; 4.P.169.239; 4.P.169.154;
4.P.169.157; 4.P.169.166; 4.P.169.169; 4.P.169.172; 4.P.169.175; 4.P.169.240;
4.P.169.244; 4.P.172.228; 4.P.172.229; 4.P.172.230; 4.P.172.231; 4.P.172.236;
20 4.P.172.237; 4.P.172.238; 4.P.172.239; 4.P.172.154; 4.P.172.157; 4.P.172.166;
4.P.172.169; 4.P.172.172; 4.P.172.175; 4.P.172.240; 4.P.172.244; 4.P.175.228;
4.P.175.229; 4.P.175.230; 4.P.175.231; 4.P.175.236; 4.P.175.237; 4.P.175.238;
4.P.175.239; 4.P.175.154; 4.P.175.157; 4.P.175.166; 4.P.175.169; 4.P.175.172;
4.P.175.175; 4.P.175.240; 4.P.175.244; 4.P.240.228; 4.P.240.229; 4.P.240.230;
25 4.P.240.231; 4.P.240.236; 4.P.240.237; 4.P.240.238; 4.P.240.239; 4.P.240.154;
4.P.240.157; 4.P.240.166; 4.P.240.169; 4.P.240.172; 4.P.240.175; 4.P.240.240;
4.P.240.244; 4.P.244.228; 4.P.244.229; 4.P.244.230; 4.P.244.231; 4.P.244.236;
4.P.244.237; 4.P.244.238; 4.P.244.239; 4.P.244.154; 4.P.244.157; 4.P.244.166;
4.P.244.169; 4.P.244.172; 4.P.244.175; 4.P.244.240; 4.P.244.244;

Prodrugs of 4.U

- 4.U.228.228; 4.U.228.229; 4.U.228.230; 4.U.228.231; 4.U.228.236;
4.U.228.237; 4.U.228.238; 4.U.228.239; 4.U.228.154; 4.U.228.157;
4.U.228.166; 4.U.228.169; 4.U.228.172; 4.U.228.175; 4.U.228.240;
5 4.U.228.244; 4.U.229.228; 4.U.229.229; 4.U.229.230; 4.U.229.231;
4.U.229.236; 4.U.229.237; 4.U.229.238; 4.U.229.239; 4.U.229.154;
4.U.229.157; 4.U.229.166; 4.U.229.169; 4.U.229.172; 4.U.229.175;
4.U.229.240; 4.U.229.244; 4.U.230.228; 4.U.230.229; 4.U.230.230;
4.U.230.231; 4.U.230.236; 4.U.230.237; 4.U.230.238; 4.U.230.239;
10 4.U.230.154; 4.U.230.157; 4.U.230.166; 4.U.230.169; 4.U.230.172;
4.U.230.175; 4.U.230.240; 4.U.230.244; 4.U.231.228; 4.U.231.229;
4.U.231.230; 4.U.231.231; 4.U.231.236; 4.U.231.237; 4.U.231.238;
4.U.231.239; 4.U.231.154; 4.U.231.157; 4.U.231.166; 4.U.231.169;
4.U.231.172; 4.U.231.175; 4.U.231.240; 4.U.231.244; 4.U.236.228;
15 4.U.236.229; 4.U.236.230; 4.U.236.231; 4.U.236.236; 4.U.236.237;
4.U.236.238; 4.U.236.239; 4.U.236.154; 4.U.236.157; 4.U.236.166;
4.U.236.169; 4.U.236.172; 4.U.236.175; 4.U.236.240; 4.U.236.244;
4.U.237.228; 4.U.237.229; 4.U.237.230; 4.U.237.231; 4.U.237.236;
4.U.237.237; 4.U.237.238; 4.U.237.239; 4.U.237.154; 4.U.237.157;
20 4.U.237.166; 4.U.237.169; 4.U.237.172; 4.U.237.175; 4.U.237.240;
4.U.237.244; 4.U.238.228; 4.U.238.229; 4.U.238.230; 4.U.238.231;
4.U.238.236; 4.U.238.237; 4.U.238.238; 4.U.238.239; 4.U.238.154;
4.U.238.157; 4.U.238.166; 4.U.238.169; 4.U.238.172; 4.U.238.175;
4.U.238.240; 4.U.238.244; 4.U.239.228; 4.U.239.229; 4.U.239.230;
25 4.U.239.231; 4.U.239.236; 4.U.239.237; 4.U.239.238; 4.U.239.239;
4.U.239.154; 4.U.239.157; 4.U.239.166; 4.U.239.169; 4.U.239.172;
4.U.239.175; 4.U.239.240; 4.U.239.244; 4.U.154.228; 4.U.154.229;
4.U.154.230; 4.U.154.231; 4.U.154.236; 4.U.154.237; 4.U.154.238;
4.U.154.239; 4.U.154.154; 4.U.154.157; 4.U.154.166; 4.U.154.169;
30 4.U.154.172; 4.U.154.175; 4.U.154.240; 4.U.154.244; 4.U.157.228;

- 4.U.157.229; 4.U.157.230; 4.U.157.231; 4.U.157.236; 4.U.157.237;
4.U.157.238; 4.U.157.239; 4.U.157.154; 4.U.157.157; 4.U.157.166;
4.U.157.169; 4.U.157.172; 4.U.157.175; 4.U.157.240; 4.U.157.244;
4.U.166.228; 4.U.166.229; 4.U.166.230; 4.U.166.231; 4.U.166.236;
5 4.U.166.237; 4.U.166.238; 4.U.166.239; 4.U.166.154; 4.U.166.157;
4.U.166.166; 4.U.166.169; 4.U.166.172; 4.U.166.175; 4.U.166.240;
4.U.166.244; 4.U.169.228; 4.U.169.229; 4.U.169.230; 4.U.169.231;
4.U.169.236; 4.U.169.237; 4.U.169.238; 4.U.169.239; 4.U.169.154;
4.U.169.157; 4.U.169.166; 4.U.169.169; 4.U.169.172; 4.U.169.175;
10 4.U.169.240; 4.U.169.244; 4.U.172.228; 4.U.172.229; 4.U.172.230;
4.U.172.231; 4.U.172.236; 4.U.172.237; 4.U.172.238; 4.U.172.239;
4.U.172.154; 4.U.172.157; 4.U.172.166; 4.U.172.169; 4.U.172.172;
4.U.172.175; 4.U.172.240; 4.U.172.244; 4.U.175.228; 4.U.175.229;
4.U.175.230; 4.U.175.231; 4.U.175.236; 4.U.175.237; 4.U.175.238;
15 4.U.175.239; 4.U.175.154; 4.U.175.157; 4.U.175.166; 4.U.175.169;
4.U.175.172; 4.U.175.175; 4.U.175.240; 4.U.175.244; 4.U.240.228;
4.U.240.229; 4.U.240.230; 4.U.240.231; 4.U.240.236; 4.U.240.237;
4.U.240.238; 4.U.240.239; 4.U.240.154; 4.U.240.157; 4.U.240.166;
4.U.240.169; 4.U.240.172; 4.U.240.175; 4.U.240.240; 4.U.240.244;
20 4.U.244.228; 4.U.244.229; 4.U.244.230; 4.U.244.231; 4.U.244.236;
4.U.244.237; 4.U.244.238; 4.U.244.239; 4.U.244.154; 4.U.244.157;
4.U.244.166; 4.U.244.169; 4.U.244.172; 4.U.244.175; 4.U.244.240;
4.U.244.244;
- 25 Prodrugs of 4.W
4.W.228.228; 4.W.228.229; 4.W.228.230; 4.W.228.231; 4.W.228.236;
4.W.228.237; 4.W.228.238; 4.W.228.239; 4.W.228.154; 4.W.228.157;
4.W.228.166; 4.W.228.169; 4.W.228.172; 4.W.228.175; 4.W.228.240;
4.W.228.244; 4.W.229.228; 4.W.229.229; 4.W.229.230; 4.W.229.231;
30 4.W.229.236; 4.W.229.237; 4.W.229.238; 4.W.229.239; 4.W.229.154;

4.W.229.157; 4.W.229.166; 4.W.229.169; 4.W.229.172; 4.W.229.175;
4.W.229.240; 4.W.229.244; 4.W.230.228; 4.W.230.229; 4.W.230.230;
4.W.230.231; 4.W.230.236; 4.W.230.237; 4.W.230.238; 4.W.230.239;
4.W.230.154; 4.W.230.157; 4.W.230.166; 4.W.230.169; 4.W.230.172;
5 4.W.230.175; 4.W.230.240; 4.W.230.244; 4.W.231.228; 4.W.231.229;
4.W.231.230; 4.W.231.231; 4.W.231.236; 4.W.231.237; 4.W.231.238;
4.W.231.239; 4.W.231.154; 4.W.231.157; 4.W.231.166; 4.W.231.169;
4.W.231.172; 4.W.231.175; 4.W.231.240; 4.W.231.244; 4.W.236.228;
4.W.236.229; 4.W.236.230; 4.W.236.231; 4.W.236.236; 4.W.236.237;
10 4.W.236.238; 4.W.236.239; 4.W.236.154; 4.W.236.157; 4.W.236.166;
4.W.236.169; 4.W.236.172; 4.W.236.175; 4.W.236.240; 4.W.236.244;
4.W.237.228; 4.W.237.229; 4.W.237.230; 4.W.237.231; 4.W.237.236;
4.W.237.237; 4.W.237.238; 4.W.237.239; 4.W.237.154; 4.W.237.157;
4.W.237.166; 4.W.237.169; 4.W.237.172; 4.W.237.175; 4.W.237.240;
15 4.W.237.244; 4.W.238.228; 4.W.238.229; 4.W.238.230; 4.W.238.231;
4.W.238.236; 4.W.238.237; 4.W.238.238; 4.W.238.239; 4.W.238.154;
4.W.238.157; 4.W.238.166; 4.W.238.169; 4.W.238.172; 4.W.238.175;
4.W.238.240; 4.W.238.244; 4.W.239.228; 4.W.239.229; 4.W.239.230;
4.W.239.231; 4.W.239.236; 4.W.239.237; 4.W.239.238; 4.W.239.239;
20 4.W.239.154; 4.W.239.157; 4.W.239.166; 4.W.239.169; 4.W.239.172;
4.W.239.175; 4.W.239.240; 4.W.239.244; 4.W.154.228; 4.W.154.229;
4.W.154.230; 4.W.154.231; 4.W.154.236; 4.W.154.237; 4.W.154.238;
4.W.154.239; 4.W.154.154; 4.W.154.157; 4.W.154.166; 4.W.154.169;
4.W.154.172; 4.W.154.175; 4.W.154.240; 4.W.154.244; 4.W.157.228;
25 4.W.157.229; 4.W.157.230; 4.W.157.231; 4.W.157.236; 4.W.157.237;
4.W.157.238; 4.W.157.239; 4.W.157.154; 4.W.157.157; 4.W.157.166;
4.W.157.169; 4.W.157.172; 4.W.157.175; 4.W.157.240; 4.W.157.244;
4.W.166.228; 4.W.166.229; 4.W.166.230; 4.W.166.231; 4.W.166.236;
4.W.166.237; 4.W.166.238; 4.W.166.239; 4.W.166.154; 4.W.166.157;
30 4.W.166.166; 4.W.166.169; 4.W.166.172; 4.W.166.175; 4.W.166.240;

4.W.166.244; 4.W.169.228; 4.W.169.229; 4.W.169.230; 4.W.169.231;
4.W.169.236; 4.W.169.237; 4.W.169.238; 4.W.169.239; 4.W.169.154;
4.W.169.157; 4.W.169.166; 4.W.169.169; 4.W.169.172; 4.W.169.175;
4.W.169.240; 4.W.169.244; 4.W.172.228; 4.W.172.229; 4.W.172.230;
5 4.W.172.231; 4.W.172.236; 4.W.172.237; 4.W.172.238; 4.W.172.239;
4.W.172.154; 4.W.172.157; 4.W.172.166; 4.W.172.169; 4.W.172.172;
4.W.172.175; 4.W.172.240; 4.W.172.244; 4.W.175.228; 4.W.175.229;
4.W.175.230; 4.W.175.231; 4.W.175.236; 4.W.175.237; 4.W.175.238;
4.W.175.239; 4.W.175.154; 4.W.175.157; 4.W.175.166; 4.W.175.169;
10 4.W.175.172; 4.W.175.175; 4.W.175.240; 4.W.175.244; 4.W.240.228;
4.W.240.229; 4.W.240.230; 4.W.240.231; 4.W.240.236; 4.W.240.237;
4.W.240.238; 4.W.240.239; 4.W.240.154; 4.W.240.157; 4.W.240.166;
4.W.240.169; 4.W.240.172; 4.W.240.175; 4.W.240.240; 4.W.240.244;
4.W.244.228; 4.W.244.229; 4.W.244.230; 4.W.244.231; 4.W.244.236;
15 4.W.244.237; 4.W.244.238; 4.W.244.239; 4.W.244.154; 4.W.244.157;
4.W.244.166; 4.W.244.169; 4.W.244.172; 4.W.244.175; 4.W.244.240;
4.W.244.244;

Prodrugs of 4.Y

20 4.Y.228.228; 4.Y.228.229; 4.Y.228.230; 4.Y.228.231; 4.Y.228.236;
4.Y.228.237; 4.Y.228.238; 4.Y.228.239; 4.Y.228.154; 4.Y.228.157; 4.Y.228.166;
4.Y.228.169; 4.Y.228.172; 4.Y.228.175; 4.Y.228.240; 4.Y.228.244; 4.Y.229.228;
4.Y.229.229; 4.Y.229.230; 4.Y.229.231; 4.Y.229.236; 4.Y.229.237; 4.Y.229.238;
4.Y.229.239; 4.Y.229.154; 4.Y.229.157; 4.Y.229.166; 4.Y.229.169; 4.Y.229.172;
25 4.Y.229.175; 4.Y.229.240; 4.Y.229.244; 4.Y.230.228; 4.Y.230.229; 4.Y.230.230;
4.Y.230.231; 4.Y.230.236; 4.Y.230.237; 4.Y.230.238; 4.Y.230.239; 4.Y.230.154;
4.Y.230.157; 4.Y.230.166; 4.Y.230.169; 4.Y.230.172; 4.Y.230.175; 4.Y.230.240;
4.Y.230.244; 4.Y.231.228; 4.Y.231.229; 4.Y.231.230; 4.Y.231.231; 4.Y.231.236;
4.Y.231.237; 4.Y.231.238; 4.Y.231.239; 4.Y.231.154; 4.Y.231.157; 4.Y.231.166;
30 4.Y.231.169; 4.Y.231.172; 4.Y.231.175; 4.Y.231.240; 4.Y.231.244; 4.Y.236.228;

4.Y.236.229; 4.Y.236.230; 4.Y.236.231; 4.Y.236.236; 4.Y.236.237; 4.Y.236.238;
4.Y.236.239; 4.Y.236.154; 4.Y.236.157; 4.Y.236.166; 4.Y.236.169; 4.Y.236.172;
4.Y.236.175; 4.Y.236.240; 4.Y.236.244; 4.Y.237.228; 4.Y.237.229; 4.Y.237.230;
4.Y.237.231; 4.Y.237.236; 4.Y.237.237; 4.Y.237.238; 4.Y.237.239; 4.Y.237.154;
5 4.Y.237.157; 4.Y.237.166; 4.Y.237.169; 4.Y.237.172; 4.Y.237.175; 4.Y.237.240;
4.Y.237.244; 4.Y.238.228; 4.Y.238.229; 4.Y.238.230; 4.Y.238.231; 4.Y.238.236;
4.Y.238.237; 4.Y.238.238; 4.Y.238.239; 4.Y.238.154; 4.Y.238.157; 4.Y.238.166;
4.Y.238.169; 4.Y.238.172; 4.Y.238.175; 4.Y.238.240; 4.Y.238.244; 4.Y.239.228;
4.Y.239.229; 4.Y.239.230; 4.Y.239.231; 4.Y.239.236; 4.Y.239.237; 4.Y.239.238;
10 4.Y.239.239; 4.Y.239.154; 4.Y.239.157; 4.Y.239.166; 4.Y.239.169; 4.Y.239.172;
4.Y.239.175; 4.Y.239.240; 4.Y.239.244; 4.Y.154.228; 4.Y.154.229; 4.Y.154.230;
4.Y.154.231; 4.Y.154.236; 4.Y.154.237; 4.Y.154.238; 4.Y.154.239; 4.Y.154.154;
4.Y.154.157; 4.Y.154.166; 4.Y.154.169; 4.Y.154.172; 4.Y.154.175; 4.Y.154.240;
4.Y.154.244; 4.Y.157.228; 4.Y.157.229; 4.Y.157.230; 4.Y.157.231; 4.Y.157.236;
15 4.Y.157.237; 4.Y.157.238; 4.Y.157.239; 4.Y.157.154; 4.Y.157.157; 4.Y.157.166;
4.Y.157.169; 4.Y.157.172; 4.Y.157.175; 4.Y.157.240; 4.Y.157.244; 4.Y.166.228;
4.Y.166.229; 4.Y.166.230; 4.Y.166.231; 4.Y.166.236; 4.Y.166.237; 4.Y.166.238;
4.Y.166.239; 4.Y.166.154; 4.Y.166.157; 4.Y.166.166; 4.Y.166.169; 4.Y.166.172;
4.Y.166.175; 4.Y.166.240; 4.Y.166.244; 4.Y.169.228; 4.Y.169.229; 4.Y.169.230;
20 4.Y.169.231; 4.Y.169.236; 4.Y.169.237; 4.Y.169.238; 4.Y.169.239; 4.Y.169.154;
4.Y.169.157; 4.Y.169.166; 4.Y.169.169; 4.Y.169.172; 4.Y.169.175; 4.Y.169.240;
4.Y.169.244; 4.Y.172.228; 4.Y.172.229; 4.Y.172.230; 4.Y.172.231; 4.Y.172.236;
4.Y.172.237; 4.Y.172.238; 4.Y.172.239; 4.Y.172.154; 4.Y.172.157; 4.Y.172.166;
4.Y.172.169; 4.Y.172.172; 4.Y.172.175; 4.Y.172.240; 4.Y.172.244; 4.Y.175.228;
25 4.Y.175.229; 4.Y.175.230; 4.Y.175.231; 4.Y.175.236; 4.Y.175.237; 4.Y.175.238;
4.Y.175.239; 4.Y.175.154; 4.Y.175.157; 4.Y.175.166; 4.Y.175.169; 4.Y.175.172;
4.Y.175.175; 4.Y.175.240; 4.Y.175.244; 4.Y.240.228; 4.Y.240.229; 4.Y.240.230;
4.Y.240.231; 4.Y.240.236; 4.Y.240.237; 4.Y.240.238; 4.Y.240.239; 4.Y.240.154;
4.Y.240.157; 4.Y.240.166; 4.Y.240.169; 4.Y.240.172; 4.Y.240.175; 4.Y.240.240;
30 4.Y.240.244; 4.Y.244.228; 4.Y.244.229; 4.Y.244.230; 4.Y.244.231; 4.Y.244.236;

4.Y.244.237; 4.Y.244.238; 4.Y.244.239; 4.Y.244.154; 4.Y.244.157; 4.Y.244.166;
4.Y.244.169; 4.Y.244.172; 4.Y.244.175; 4.Y.244.240; 4.Y.244.244;

Prodrugs of 5.B

- 5 5.B.228.228; 5.B.228.229; 5.B.228.230; 5.B.228.231; 5.B.228.236;
5.B.228.237; 5.B.228.238; 5.B.228.239; 5.B.228.154; 5.B.228.157; 5.B.228.166;
5.B.228.169; 5.B.228.172; 5.B.228.175; 5.B.228.240; 5.B.228.244; 5.B.229.228;
5.B.229.229; 5.B.229.230; 5.B.229.231; 5.B.229.236; 5.B.229.237; 5.B.229.238;
5.B.229.239; 5.B.229.154; 5.B.229.157; 5.B.229.166; 5.B.229.169; 5.B.229.172;
10 5.B.229.175; 5.B.229.240; 5.B.229.244; 5.B.230.228; 5.B.230.229; 5.B.230.230;
5.B.230.231; 5.B.230.236; 5.B.230.237; 5.B.230.238; 5.B.230.239; 5.B.230.154;
5.B.230.157; 5.B.230.166; 5.B.230.169; 5.B.230.172; 5.B.230.175; 5.B.230.240;
5.B.230.244; 5.B.231.228; 5.B.231.229; 5.B.231.230; 5.B.231.231; 5.B.231.236;
5.B.231.237; 5.B.231.238; 5.B.231.239; 5.B.231.154; 5.B.231.157; 5.B.231.166;
15 5.B.231.169; 5.B.231.172; 5.B.231.175; 5.B.231.240; 5.B.231.244; 5.B.236.228;
5.B.236.229; 5.B.236.230; 5.B.236.231; 5.B.236.236; 5.B.236.237; 5.B.236.238;
5.B.236.239; 5.B.236.154; 5.B.236.157; 5.B.236.166; 5.B.236.169; 5.B.236.172;
5.B.236.175; 5.B.236.240; 5.B.236.244; 5.B.237.228; 5.B.237.229; 5.B.237.230;
5.B.237.231; 5.B.237.236; 5.B.237.237; 5.B.237.238; 5.B.237.239; 5.B.237.154;
20 5.B.237.157; 5.B.237.166; 5.B.237.169; 5.B.237.172; 5.B.237.175; 5.B.237.240;
5.B.237.244; 5.B.238.228; 5.B.238.229; 5.B.238.230; 5.B.238.231; 5.B.238.236;
5.B.238.237; 5.B.238.238; 5.B.238.239; 5.B.238.154; 5.B.238.157; 5.B.238.166;
5.B.238.169; 5.B.238.172; 5.B.238.175; 5.B.238.240; 5.B.238.244; 5.B.239.228;
5.B.239.229; 5.B.239.230; 5.B.239.231; 5.B.239.236; 5.B.239.237; 5.B.239.238;
25 5.B.239.239; 5.B.239.154; 5.B.239.157; 5.B.239.166; 5.B.239.169; 5.B.239.172;
5.B.239.175; 5.B.239.240; 5.B.239.244; 5.B.154.228; 5.B.154.229; 5.B.154.230;
5.B.154.231; 5.B.154.236; 5.B.154.237; 5.B.154.238; 5.B.154.239; 5.B.154.154;
5.B.154.157; 5.B.154.166; 5.B.154.169; 5.B.154.172; 5.B.154.175; 5.B.154.240;
5.B.154.244; 5.B.157.228; 5.B.157.229; 5.B.157.230; 5.B.157.231; 5.B.157.236;
30 5.B.157.237; 5.B.157.238; 5.B.157.239; 5.B.157.154; 5.B.157.157; 5.B.157.166;

5.B.157.169; 5.B.157.172; 5.B.157.175; 5.B.157.240; 5.B.157.244; 5.B.166.228;
5.B.166.229; 5.B.166.230; 5.B.166.231; 5.B.166.236; 5.B.166.237; 5.B.166.238;
5.B.166.239; 5.B.166.154; 5.B.166.157; 5.B.166.166; 5.B.166.169; 5.B.166.172;
5.B.166.175; 5.B.166.240; 5.B.166.244; 5.B.169.228; 5.B.169.229; 5.B.169.230;
5 5.B.169.231; 5.B.169.236; 5.B.169.237; 5.B.169.238; 5.B.169.239; 5.B.169.154;
5.B.169.157; 5.B.169.166; 5.B.169.169; 5.B.169.172; 5.B.169.175; 5.B.169.240;
5.B.169.244; 5.B.172.228; 5.B.172.229; 5.B.172.230; 5.B.172.231; 5.B.172.236;
5.B.172.237; 5.B.172.238; 5.B.172.239; 5.B.172.154; 5.B.172.157; 5.B.172.166;
5.B.172.169; 5.B.172.172; 5.B.172.175; 5.B.172.240; 5.B.172.244; 5.B.175.228;
10 5.B.175.229; 5.B.175.230; 5.B.175.231; 5.B.175.236; 5.B.175.237; 5.B.175.238;
5.B.175.239; 5.B.175.154; 5.B.175.157; 5.B.175.166; 5.B.175.169; 5.B.175.172;
5.B.175.175; 5.B.175.240; 5.B.175.244; 5.B.240.228; 5.B.240.229; 5.B.240.230;
5.B.240.231; 5.B.240.236; 5.B.240.237; 5.B.240.238; 5.B.240.239; 5.B.240.154;
5.B.240.157; 5.B.240.166; 5.B.240.169; 5.B.240.172; 5.B.240.175; 5.B.240.240;
15 5.B.240.244; 5.B.244.228; 5.B.244.229; 5.B.244.230; 5.B.244.231; 5.B.244.236;
5.B.244.237; 5.B.244.238; 5.B.244.239; 5.B.244.154; 5.B.244.157; 5.B.244.166;
5.B.244.169; 5.B.244.172; 5.B.244.175; 5.B.244.240; 5.B.244.244;

Prodrugs of 5.D

20 5.D.228.228; 5.D.228.229; 5.D.228.230; 5.D.228.231; 5.D.228.236;
5.D.228.237; 5.D.228.238; 5.D.228.239; 5.D.228.154; 5.D.228.157;
5.D.228.166; 5.D.228.169; 5.D.228.172; 5.D.228.175; 5.D.228.240;
5.D.228.244; 5.D.229.228; 5.D.229.229; 5.D.229.230; 5.D.229.231;
5.D.229.236; 5.D.229.237; 5.D.229.238; 5.D.229.239; 5.D.229.154;
25 5.D.229.157; 5.D.229.166; 5.D.229.169; 5.D.229.172; 5.D.229.175;
5.D.229.240; 5.D.229.244; 5.D.230.228; 5.D.230.229; 5.D.230.230;
5.D.230.231; 5.D.230.236; 5.D.230.237; 5.D.230.238; 5.D.230.239;
5.D.230.154; 5.D.230.157; 5.D.230.166; 5.D.230.169; 5.D.230.172;
5.D.230.175; 5.D.230.240; 5.D.230.244; 5.D.231.228; 5.D.231.229;
30 5.D.231.230; 5.D.231.231; 5.D.231.236; 5.D.231.237; 5.D.231.238;

5.D.231.239; 5.D.231.154; 5.D.231.157; 5.D.231.166; 5.D.231.169;
5.D.231.172; 5.D.231.175; 5.D.231.240; 5.D.231.244; 5.D.236.228;
5.D.236.229; 5.D.236.230; 5.D.236.231; 5.D.236.236; 5.D.236.237;
5.D.236.238; 5.D.236.239; 5.D.236.154; 5.D.236.157; 5.D.236.166;
5 5.D.236.169; 5.D.236.172; 5.D.236.175; 5.D.236.240; 5.D.236.244;
5.D.237.228; 5.D.237.229; 5.D.237.230; 5.D.237.231; 5.D.237.236;
5.D.237.237; 5.D.237.238; 5.D.237.239; 5.D.237.154; 5.D.237.157;
5.D.237.166; 5.D.237.169; 5.D.237.172; 5.D.237.175; 5.D.237.240;
5.D.237.244; 5.D.238.228; 5.D.238.229; 5.D.238.230; 5.D.238.231;
10 5.D.238.236; 5.D.238.237; 5.D.238.238; 5.D.238.239; 5.D.238.154;
5.D.238.157; 5.D.238.166; 5.D.238.169; 5.D.238.172; 5.D.238.175;
5.D.238.240; 5.D.238.244; 5.D.239.228; 5.D.239.229; 5.D.239.230;
5.D.239.231; 5.D.239.236; 5.D.239.237; 5.D.239.238; 5.D.239.239;
5.D.239.154; 5.D.239.157; 5.D.239.166; 5.D.239.169; 5.D.239.172;
15 5.D.239.175; 5.D.239.240; 5.D.239.244; 5.D.154.228; 5.D.154.229;
5.D.154.230; 5.D.154.231; 5.D.154.236; 5.D.154.237; 5.D.154.238;
5.D.154.239; 5.D.154.154; 5.D.154.157; 5.D.154.166; 5.D.154.169;
5.D.154.172; 5.D.154.175; 5.D.154.240; 5.D.154.244; 5.D.157.228;
5.D.157.229; 5.D.157.230; 5.D.157.231; 5.D.157.236; 5.D.157.237;
20 5.D.157.238; 5.D.157.239; 5.D.157.154; 5.D.157.157; 5.D.157.166;
5.D.157.169; 5.D.157.172; 5.D.157.175; 5.D.157.240; 5.D.157.244;
5.D.166.228; 5.D.166.229; 5.D.166.230; 5.D.166.231; 5.D.166.236;
5.D.166.237; 5.D.166.238; 5.D.166.239; 5.D.166.154; 5.D.166.157;
5.D.166.166; 5.D.166.169; 5.D.166.172; 5.D.166.175; 5.D.166.240;
25 5.D.166.244; 5.D.169.228; 5.D.169.229; 5.D.169.230; 5.D.169.231;
5.D.169.236; 5.D.169.237; 5.D.169.238; 5.D.169.239; 5.D.169.154;
5.D.169.157; 5.D.169.166; 5.D.169.169; 5.D.169.172; 5.D.169.175;
5.D.169.240; 5.D.169.244; 5.D.172.228; 5.D.172.229; 5.D.172.230;
5.D.172.231; 5.D.172.236; 5.D.172.237; 5.D.172.238; 5.D.172.239;
30 5.D.172.154; 5.D.172.157; 5.D.172.166; 5.D.172.169; 5.D.172.172;

5.D.172.175; 5.D.172.240; 5.D.172.244; 5.D.175.228; 5.D.175.229;
 5.D.175.230; 5.D.175.231; 5.D.175.236; 5.D.175.237; 5.D.175.238;
 5.D.175.239; 5.D.175.154; 5.D.175.157; 5.D.175.166; 5.D.175.169;
 5.D.175.172; 5.D.175.175; 5.D.175.240; 5.D.175.244; 5.D.240.228;
 5 5.D.240.229; 5.D.240.230; 5.D.240.231; 5.D.240.236; 5.D.240.237;
 5.D.240.238; 5.D.240.239; 5.D.240.154; 5.D.240.157; 5.D.240.166;
 5.D.240.169; 5.D.240.172; 5.D.240.175; 5.D.240.240; 5.D.240.244;
 5.D.244.228; 5.D.244.229; 5.D.244.230; 5.D.244.231; 5.D.244.236;
 5.D.244.237; 5.D.244.238; 5.D.244.239; 5.D.244.154; 5.D.244.157;
 10 5.D.244.166; 5.D.244.169; 5.D.244.172; 5.D.244.175; 5.D.244.240;
 5.D.244.244;

Prodrugs of 5.E

5.E.228.228; 5.E.228.229; 5.E.228.230; 5.E.228.231; 5.E.228.236;
 15 5.E.228.237; 5.E.228.238; 5.E.228.239; 5.E.228.154; 5.E.228.157; 5.E.228.166;
 5.E.228.169; 5.E.228.172; 5.E.228.175; 5.E.228.240; 5.E.228.244; 5.E.229.228;
 5.E.229.229; 5.E.229.230; 5.E.229.231; 5.E.229.236; 5.E.229.237; 5.E.229.238;
 5.E.229.239; 5.E.229.154; 5.E.229.157; 5.E.229.166; 5.E.229.169; 5.E.229.172;
 5.E.229.175; 5.E.229.240; 5.E.229.244; 5.E.230.228; 5.E.230.229; 5.E.230.230;
 20 5.E.230.231; 5.E.230.236; 5.E.230.237; 5.E.230.238; 5.E.230.239; 5.E.230.154;
 5.E.230.157; 5.E.230.166; 5.E.230.169; 5.E.230.172; 5.E.230.175; 5.E.230.240;
 5.E.230.244; 5.E.231.228; 5.E.231.229; 5.E.231.230; 5.E.231.231; 5.E.231.236;
 5.E.231.237; 5.E.231.238; 5.E.231.239; 5.E.231.154; 5.E.231.157; 5.E.231.166;
 5.E.231.169; 5.E.231.172; 5.E.231.175; 5.E.231.240; 5.E.231.244; 5.E.236.228;
 25 5.E.236.229; 5.E.236.230; 5.E.236.231; 5.E.236.236; 5.E.236.237; 5.E.236.238;
 5.E.236.239; 5.E.236.154; 5.E.236.157; 5.E.236.166; 5.E.236.169; 5.E.236.172;
 5.E.236.175; 5.E.236.240; 5.E.236.244; 5.E.237.228; 5.E.237.229; 5.E.237.230;
 5.E.237.231; 5.E.237.236; 5.E.237.237; 5.E.237.238; 5.E.237.239; 5.E.237.154;
 5.E.237.157; 5.E.237.166; 5.E.237.169; 5.E.237.172; 5.E.237.175; 5.E.237.240;
 30 5.E.237.244; 5.E.238.228; 5.E.238.229; 5.E.238.230; 5.E.238.231; 5.E.238.236;

5.E.238.237; 5.E.238.238; 5.E.238.239; 5.E.238.154; 5.E.238.157; 5.E.238.166;
5.E.238.169; 5.E.238.172; 5.E.238.175; 5.E.238.240; 5.E.238.244; 5.E.239.228;
5.E.239.229; 5.E.239.230; 5.E.239.231; 5.E.239.236; 5.E.239.237; 5.E.239.238;
5.E.239.239; 5.E.239.154; 5.E.239.157; 5.E.239.166; 5.E.239.169; 5.E.239.172;
5 5.E.239.175; 5.E.239.240; 5.E.239.244; 5.E.154.228; 5.E.154.229; 5.E.154.230;
5.E.154.231; 5.E.154.236; 5.E.154.237; 5.E.154.238; 5.E.154.239; 5.E.154.154;
5.E.154.157; 5.E.154.166; 5.E.154.169; 5.E.154.172; 5.E.154.175; 5.E.154.240;
5.E.154.244; 5.E.157.228; 5.E.157.229; 5.E.157.230; 5.E.157.231; 5.E.157.236;
5.E.157.237; 5.E.157.238; 5.E.157.239; 5.E.157.154; 5.E.157.157; 5.E.157.166;
10 5.E.157.169; 5.E.157.172; 5.E.157.175; 5.E.157.240; 5.E.157.244; 5.E.166.228;
5.E.166.229; 5.E.166.230; 5.E.166.231; 5.E.166.236; 5.E.166.237; 5.E.166.238;
5.E.166.239; 5.E.166.154; 5.E.166.157; 5.E.166.166; 5.E.166.169; 5.E.166.172;
5.E.166.175; 5.E.166.240; 5.E.166.244; 5.E.169.228; 5.E.169.229; 5.E.169.230;
5.E.169.231; 5.E.169.236; 5.E.169.237; 5.E.169.238; 5.E.169.239; 5.E.169.154;
15 5.E.169.157; 5.E.169.166; 5.E.169.169; 5.E.169.172; 5.E.169.175; 5.E.169.240;
5.E.169.244; 5.E.172.228; 5.E.172.229; 5.E.172.230; 5.E.172.231; 5.E.172.236;
5.E.172.237; 5.E.172.238; 5.E.172.239; 5.E.172.154; 5.E.172.157; 5.E.172.166;
5.E.172.169; 5.E.172.172; 5.E.172.175; 5.E.172.240; 5.E.172.244; 5.E.175.228;
5.E.175.229; 5.E.175.230; 5.E.175.231; 5.E.175.236; 5.E.175.237; 5.E.175.238;
20 5.E.175.239; 5.E.175.154; 5.E.175.157; 5.E.175.166; 5.E.175.169; 5.E.175.172;
5.E.175.175; 5.E.175.240; 5.E.175.244; 5.E.240.228; 5.E.240.229; 5.E.240.230;
5.E.240.231; 5.E.240.236; 5.E.240.237; 5.E.240.238; 5.E.240.239; 5.E.240.154;
5.E.240.157; 5.E.240.166; 5.E.240.169; 5.E.240.172; 5.E.240.175; 5.E.240.240;
5.E.240.244; 5.E.244.228; 5.E.244.229; 5.E.244.230; 5.E.244.231; 5.E.244.236;
25 5.E.244.237; 5.E.244.238; 5.E.244.239; 5.E.244.154; 5.E.244.157; 5.E.244.166;
5.E.244.169; 5.E.244.172; 5.E.244.175; 5.E.244.240; 5.E.244.244;

Prodrugs of 5.G

5.G.228.228; 5.G.228.229; 5.G.228.230; 5.G.228.231; 5.G.228.236;
30 5.G.228.237; 5.G.228.238; 5.G.228.239; 5.G.228.154; 5.G.228.157;

5.G.228.166; 5.G.228.169; 5.G.228.172; 5.G.228.175; 5.G.228.240;
5.G.228.244; 5.G.229.228; 5.G.229.229; 5.G.229.230; 5.G.229.231;
5.G.229.236; 5.G.229.237; 5.G.229.238; 5.G.229.239; 5.G.229.154;
5.G.229.157; 5.G.229.166; 5.G.229.169; 5.G.229.172; 5.G.229.175;
5 5.G.229.240; 5.G.229.244; 5.G.230.228; 5.G.230.229; 5.G.230.230;
5.G.230.231; 5.G.230.236; 5.G.230.237; 5.G.230.238; 5.G.230.239;
5.G.230.154; 5.G.230.157; 5.G.230.166; 5.G.230.169; 5.G.230.172;
5.G.230.175; 5.G.230.240; 5.G.230.244; 5.G.231.228; 5.G.231.229;
5.G.231.230; 5.G.231.231; 5.G.231.236; 5.G.231.237; 5.G.231.238;
10 5.G.231.239; 5.G.231.154; 5.G.231.157; 5.G.231.166; 5.G.231.169;
5.G.231.172; 5.G.231.175; 5.G.231.240; 5.G.231.244; 5.G.236.228;
5.G.236.229; 5.G.236.230; 5.G.236.231; 5.G.236.236; 5.G.236.237;
5.G.236.238; 5.G.236.239; 5.G.236.154; 5.G.236.157; 5.G.236.166;
5.G.236.169; 5.G.236.172; 5.G.236.175; 5.G.236.240; 5.G.236.244;
15 5.G.237.228; 5.G.237.229; 5.G.237.230; 5.G.237.231; 5.G.237.236;
5.G.237.237; 5.G.237.238; 5.G.237.239; 5.G.237.154; 5.G.237.157;
5.G.237.166; 5.G.237.169; 5.G.237.172; 5.G.237.175; 5.G.237.240;
5.G.237.244; 5.G.238.228; 5.G.238.229; 5.G.238.230; 5.G.238.231;
5.G.238.236; 5.G.238.237; 5.G.238.238; 5.G.238.239; 5.G.238.154;
20 5.G.238.157; 5.G.238.166; 5.G.238.169; 5.G.238.172; 5.G.238.175;
5.G.238.240; 5.G.238.244; 5.G.239.228; 5.G.239.229; 5.G.239.230;
5.G.239.231; 5.G.239.236; 5.G.239.237; 5.G.239.238; 5.G.239.239;
5.G.239.154; 5.G.239.157; 5.G.239.166; 5.G.239.169; 5.G.239.172;
5.G.239.175; 5.G.239.240; 5.G.239.244; 5.G.154.228; 5.G.154.229;
25 5.G.154.230; 5.G.154.231; 5.G.154.236; 5.G.154.237; 5.G.154.238;
5.G.154.239; 5.G.154.154; 5.G.154.157; 5.G.154.166; 5.G.154.169;
5.G.154.172; 5.G.154.175; 5.G.154.240; 5.G.154.244; 5.G.157.228;
5.G.157.229; 5.G.157.230; 5.G.157.231; 5.G.157.236; 5.G.157.237;
5.G.157.238; 5.G.157.239; 5.G.157.154; 5.G.157.157; 5.G.157.166;
30 5.G.157.169; 5.G.157.172; 5.G.157.175; 5.G.157.240; 5.G.157.244;

5.G.166.228; 5.G.166.229; 5.G.166.230; 5.G.166.231; 5.G.166.236;
 5.G.166.237; 5.G.166.238; 5.G.166.239; 5.G.166.154; 5.G.166.157;
 5.G.166.166; 5.G.166.169; 5.G.166.172; 5.G.166.175; 5.G.166.240;
 5.G.166.244; 5.G.169.228; 5.G.169.229; 5.G.169.230; 5.G.169.231;
 5 5.G.169.236; 5.G.169.237; 5.G.169.238; 5.G.169.239; 5.G.169.154;
 5.G.169.157; 5.G.169.166; 5.G.169.169; 5.G.169.172; 5.G.169.175;
 5.G.169.240; 5.G.169.244; 5.G.172.228; 5.G.172.229; 5.G.172.230;
 5.G.172.231; 5.G.172.236; 5.G.172.237; 5.G.172.238; 5.G.172.239;
 5.G.172.154; 5.G.172.157; 5.G.172.166; 5.G.172.169; 5.G.172.172;
 10 5.G.172.175; 5.G.172.240; 5.G.172.244; 5.G.175.228; 5.G.175.229;
 5.G.175.230; 5.G.175.231; 5.G.175.236; 5.G.175.237; 5.G.175.238;
 5.G.175.239; 5.G.175.154; 5.G.175.157; 5.G.175.166; 5.G.175.169;
 5.G.175.172; 5.G.175.175; 5.G.175.240; 5.G.175.244; 5.G.240.228;
 5.G.240.229; 5.G.240.230; 5.G.240.231; 5.G.240.236; 5.G.240.237;
 15 5.G.240.238; 5.G.240.239; 5.G.240.154; 5.G.240.157; 5.G.240.166;
 5.G.240.169; 5.G.240.172; 5.G.240.175; 5.G.240.240; 5.G.240.244;
 5.G.244.228; 5.G.244.229; 5.G.244.230; 5.G.244.231; 5.G.244.236;
 5.G.244.237; 5.G.244.238; 5.G.244.239; 5.G.244.154; 5.G.244.157;
 5.G.244.166; 5.G.244.169; 5.G.244.172; 5.G.244.175; 5.G.244.240;
 20 5.G.244.244;

Prodrugs of 5.I

5.I.228.228; 5.I.228.229; 5.I.228.230; 5.I.228.231; 5.I.228.236; 5.I.228.237;
 5.I.228.238; 5.I.228.239; 5.I.228.154; 5.I.228.157; 5.I.228.166; 5.I.228.169;
 25 5.I.228.172; 5.I.228.175; 5.I.228.240; 5.I.228.244; 5.I.229.228; 5.I.229.229;
 5.I.229.230; 5.I.229.231; 5.I.229.236; 5.I.229.237; 5.I.229.238; 5.I.229.239;
 5.I.229.154; 5.I.229.157; 5.I.229.166; 5.I.229.169; 5.I.229.172; 5.I.229.175;
 5.I.229.240; 5.I.229.244; 5.I.230.228; 5.I.230.229; 5.I.230.230; 5.I.230.231;
 5.I.230.236; 5.I.230.237; 5.I.230.238; 5.I.230.239; 5.I.230.154; 5.I.230.157;
 30 5.I.230.166; 5.I.230.169; 5.I.230.172; 5.I.230.175; 5.I.230.240; 5.I.230.244;

5.I.231.228; 5.I.231.229; 5.I.231.230; 5.I.231.231; 5.I.231.236; 5.I.231.237;
5.I.231.238; 5.I.231.239; 5.I.231.154; 5.I.231.157; 5.I.231.166; 5.I.231.169;
5.I.231.172; 5.I.231.175; 5.I.231.240; 5.I.231.244; 5.I.236.228; 5.I.236.229;
5.I.236.230; 5.I.236.231; 5.I.236.236; 5.I.236.237; 5.I.236.238; 5.I.236.239;
5 5.I.236.154; 5.I.236.157; 5.I.236.166; 5.I.236.169; 5.I.236.172; 5.I.236.175;
5.I.236.240; 5.I.236.244; 5.I.237.228; 5.I.237.229; 5.I.237.230; 5.I.237.231;
5.I.237.236; 5.I.237.237; 5.I.237.238; 5.I.237.239; 5.I.237.154; 5.I.237.157;
5.I.237.166; 5.I.237.169; 5.I.237.172; 5.I.237.175; 5.I.237.240; 5.I.237.244;
5.I.238.228; 5.I.238.229; 5.I.238.230; 5.I.238.231; 5.I.238.236; 5.I.238.237;
10 5.I.238.238; 5.I.238.239; 5.I.238.154; 5.I.238.157; 5.I.238.166; 5.I.238.169;
5.I.238.172; 5.I.238.175; 5.I.238.240; 5.I.238.244; 5.I.239.228; 5.I.239.229;
5.I.239.230; 5.I.239.231; 5.I.239.236; 5.I.239.237; 5.I.239.238; 5.I.239.239;
5.I.239.154; 5.I.239.157; 5.I.239.166; 5.I.239.169; 5.I.239.172; 5.I.239.175;
5.I.239.240; 5.I.239.244; 5.I.154.228; 5.I.154.229; 5.I.154.230; 5.I.154.231;
15 5.I.154.236; 5.I.154.237; 5.I.154.238; 5.I.154.239; 5.I.154.154; 5.I.154.157;
5.I.154.166; 5.I.154.169; 5.I.154.172; 5.I.154.175; 5.I.154.240; 5.I.154.244;
5.I.157.228; 5.I.157.229; 5.I.157.230; 5.I.157.231; 5.I.157.236; 5.I.157.237;
5.I.157.238; 5.I.157.239; 5.I.157.154; 5.I.157.157; 5.I.157.166; 5.I.157.169;
5.I.157.172; 5.I.157.175; 5.I.157.240; 5.I.157.244; 5.I.166.228; 5.I.166.229;
20 5.I.166.230; 5.I.166.231; 5.I.166.236; 5.I.166.237; 5.I.166.238; 5.I.166.239;
5.I.166.154; 5.I.166.157; 5.I.166.166; 5.I.166.169; 5.I.166.172; 5.I.166.175;
5.I.166.240; 5.I.166.244; 5.I.169.228; 5.I.169.229; 5.I.169.230; 5.I.169.231;
5.I.169.236; 5.I.169.237; 5.I.169.238; 5.I.169.239; 5.I.169.154; 5.I.169.157;
5.I.169.166; 5.I.169.169; 5.I.169.172; 5.I.169.175; 5.I.169.240; 5.I.169.244;
25 5.I.172.228; 5.I.172.229; 5.I.172.230; 5.I.172.231; 5.I.172.236; 5.I.172.237;
5.I.172.238; 5.I.172.239; 5.I.172.154; 5.I.172.157; 5.I.172.166; 5.I.172.169;
5.I.172.172; 5.I.172.175; 5.I.172.240; 5.I.172.244; 5.I.175.228; 5.I.175.229;
5.I.175.230; 5.I.175.231; 5.I.175.236; 5.I.175.237; 5.I.175.238; 5.I.175.239;
5.I.175.154; 5.I.175.157; 5.I.175.166; 5.I.175.169; 5.I.175.172; 5.I.175.175;
30 5.I.175.240; 5.I.175.244; 5.I.240.228; 5.I.240.229; 5.I.240.230; 5.I.240.231;

5.I.240.236; 5.I.240.237; 5.I.240.238; 5.I.240.239; 5.I.240.154; 5.I.240.157;
5.I.240.166; 5.I.240.169; 5.I.240.172; 5.I.240.175; 5.I.240.240; 5.I.240.244;
5.I.244.228; 5.I.244.229; 5.I.244.230; 5.I.244.231; 5.I.244.236; 5.I.244.237;
5.I.244.238; 5.I.244.239; 5.I.244.154; 5.I.244.157; 5.I.244.166; 5.I.244.169;
5 5.I.244.172; 5.I.244.175; 5.I.244.240; 5.I.244.244;

Prodrugs of 5.I

5.J.228.228; 5.J.228.229; 5.J.228.230; 5.J.228.231; 5.J.228.236; 5.J.228.237;
5.J.228.238; 5.J.228.239; 5.J.228.154; 5.J.228.157; 5.J.228.166; 5.J.228.169;
10 5.J.228.172; 5.J.228.175; 5.J.228.240; 5.J.228.244; 5.J.229.228; 5.J.229.229;
5.J.229.230; 5.J.229.231; 5.J.229.236; 5.J.229.237; 5.J.229.238; 5.J.229.239;
5.J.229.154; 5.J.229.157; 5.J.229.166; 5.J.229.169; 5.J.229.172; 5.J.229.175;
5.J.229.240; 5.J.229.244; 5.J.230.228; 5.J.230.229; 5.J.230.230; 5.J.230.231;
5.J.230.236; 5.J.230.237; 5.J.230.238; 5.J.230.239; 5.J.230.154; 5.J.230.157;
15 5.J.230.166; 5.J.230.169; 5.J.230.172; 5.J.230.175; 5.J.230.240; 5.J.230.244;
5.J.231.228; 5.J.231.229; 5.J.231.230; 5.J.231.231; 5.J.231.236; 5.J.231.237;
5.J.231.238; 5.J.231.239; 5.J.231.154; 5.J.231.157; 5.J.231.166; 5.J.231.169;
5.J.231.172; 5.J.231.175; 5.J.231.240; 5.J.231.244; 5.J.236.228; 5.J.236.229;
5.J.236.230; 5.J.236.231; 5.J.236.236; 5.J.236.237; 5.J.236.238; 5.J.236.239;
20 5.J.236.154; 5.J.236.157; 5.J.236.166; 5.J.236.169; 5.J.236.172; 5.J.236.175;
5.J.236.240; 5.J.236.244; 5.J.237.228; 5.J.237.229; 5.J.237.230; 5.J.237.231;
5.J.237.236; 5.J.237.237; 5.J.237.238; 5.J.237.239; 5.J.237.154; 5.J.237.157;
5.J.237.166; 5.J.237.169; 5.J.237.172; 5.J.237.175; 5.J.237.240; 5.J.237.244;
5.J.238.228; 5.J.238.229; 5.J.238.230; 5.J.238.231; 5.J.238.236; 5.J.238.237;
25 5.J.238.238; 5.J.238.239; 5.J.238.154; 5.J.238.157; 5.J.238.166; 5.J.238.169;
5.J.238.172; 5.J.238.175; 5.J.238.240; 5.J.238.244; 5.J.239.228; 5.J.239.229;
5.J.239.230; 5.J.239.231; 5.J.239.236; 5.J.239.237; 5.J.239.238; 5.J.239.239;
5.J.239.154; 5.J.239.157; 5.J.239.166; 5.J.239.169; 5.J.239.172; 5.J.239.175;
5.J.239.240; 5.J.239.244; 5.J.154.228; 5.J.154.229; 5.J.154.230; 5.J.154.231;
30 5.J.154.236; 5.J.154.237; 5.J.154.238; 5.J.154.239; 5.J.154.154; 5.J.154.157;

5.J.154.166; 5.J.154.169; 5.J.154.172; 5.J.154.175; 5.J.154.240; 5.J.154.244;
5.J.157.228; 5.J.157.229; 5.J.157.230; 5.J.157.231; 5.J.157.236; 5.J.157.237;
5.J.157.238; 5.J.157.239; 5.J.157.154; 5.J.157.157; 5.J.157.166; 5.J.157.169;
5.J.157.172; 5.J.157.175; 5.J.157.240; 5.J.157.244; 5.J.166.228; 5.J.166.229;
5 5.J.166.230; 5.J.166.231; 5.J.166.236; 5.J.166.237; 5.J.166.238; 5.J.166.239;
5.J.166.154; 5.J.166.157; 5.J.166.166; 5.J.166.169; 5.J.166.172; 5.J.166.175;
5.J.166.240; 5.J.166.244; 5.J.169.228; 5.J.169.229; 5.J.169.230; 5.J.169.231;
5.J.169.236; 5.J.169.237; 5.J.169.238; 5.J.169.239; 5.J.169.154; 5.J.169.157;
5.J.169.166; 5.J.169.169; 5.J.169.172; 5.J.169.175; 5.J.169.240; 5.J.169.244;
10 5.J.172.228; 5.J.172.229; 5.J.172.230; 5.J.172.231; 5.J.172.236; 5.J.172.237;
5.J.172.238; 5.J.172.239; 5.J.172.154; 5.J.172.157; 5.J.172.166; 5.J.172.169;
5.J.172.172; 5.J.172.175; 5.J.172.240; 5.J.172.244; 5.J.175.228; 5.J.175.229;
5.J.175.230; 5.J.175.231; 5.J.175.236; 5.J.175.237; 5.J.175.238; 5.J.175.239;
5.J.175.154; 5.J.175.157; 5.J.175.166; 5.J.175.169; 5.J.175.172; 5.J.175.175;
15 5.J.175.240; 5.J.175.244; 5.J.240.228; 5.J.240.229; 5.J.240.230; 5.J.240.231;
5.J.240.236; 5.J.240.237; 5.J.240.238; 5.J.240.239; 5.J.240.154; 5.J.240.157;
5.J.240.166; 5.J.240.169; 5.J.240.172; 5.J.240.175; 5.J.240.240; 5.J.240.244;
5.J.244.228; 5.J.244.229; 5.J.244.230; 5.J.244.231; 5.J.244.236; 5.J.244.237;
5.J.244.238; 5.J.244.239; 5.J.244.154; 5.J.244.157; 5.J.244.166; 5.J.244.169;
20 5.J.244.172; 5.J.244.175; 5.J.244.240; 5.J.244.244;

Prodrugs of 5.L

5.L.228.228; 5.L.228.229; 5.L.228.230; 5.L.228.231; 5.L.228.236;
5.L.228.237; 5.L.228.238; 5.L.228.239; 5.L.228.154; 5.L.228.157; 5.L.228.166;
25 5.L.228.169; 5.L.228.172; 5.L.228.175; 5.L.228.240; 5.L.228.244; 5.L.229.228;
5.L.229.229; 5.L.229.230; 5.L.229.231; 5.L.229.236; 5.L.229.237; 5.L.229.238;
5.L.229.239; 5.L.229.154; 5.L.229.157; 5.L.229.166; 5.L.229.169; 5.L.229.172;
5.L.229.175; 5.L.229.240; 5.L.229.244; 5.L.230.228; 5.L.230.229; 5.L.230.230;
5.L.230.231; 5.L.230.236; 5.L.230.237; 5.L.230.238; 5.L.230.239; 5.L.230.154;
30 5.L.230.157; 5.L.230.166; 5.L.230.169; 5.L.230.172; 5.L.230.175; 5.L.230.240;

5.L.230.244; 5.L.231.228; 5.L.231.229; 5.L.231.230; 5.L.231.231; 5.L.231.236;
5.L.231.237; 5.L.231.238; 5.L.231.239; 5.L.231.154; 5.L.231.157; 5.L.231.166;
5.L.231.169; 5.L.231.172; 5.L.231.175; 5.L.231.240; 5.L.231.244; 5.L.236.228;
5.L.236.229; 5.L.236.230; 5.L.236.231; 5.L.236.236; 5.L.236.237; 5.L.236.238;
5 5.L.236.239; 5.L.236.154; 5.L.236.157; 5.L.236.166; 5.L.236.169; 5.L.236.172;
5.L.236.175; 5.L.236.240; 5.L.236.244; 5.L.237.228; 5.L.237.229; 5.L.237.230;
5.L.237.231; 5.L.237.236; 5.L.237.237; 5.L.237.238; 5.L.237.239; 5.L.237.154;
5.L.237.157; 5.L.237.166; 5.L.237.169; 5.L.237.172; 5.L.237.175; 5.L.237.240;
5.L.237.244; 5.L.238.228; 5.L.238.229; 5.L.238.230; 5.L.238.231; 5.L.238.236;
10 5.L.238.237; 5.L.238.238; 5.L.238.239; 5.L.238.154; 5.L.238.157; 5.L.238.166;
5.L.238.169; 5.L.238.172; 5.L.238.175; 5.L.238.240; 5.L.238.244; 5.L.239.228;
5.L.239.229; 5.L.239.230; 5.L.239.231; 5.L.239.236; 5.L.239.237; 5.L.239.238;
5.L.239.239; 5.L.239.154; 5.L.239.157; 5.L.239.166; 5.L.239.169; 5.L.239.172;
5.L.239.175; 5.L.239.240; 5.L.239.244; 5.L.154.228; 5.L.154.229; 5.L.154.230;
15 5.L.154.231; 5.L.154.236; 5.L.154.237; 5.L.154.238; 5.L.154.239; 5.L.154.154;
5.L.154.157; 5.L.154.166; 5.L.154.169; 5.L.154.172; 5.L.154.175; 5.L.154.240;
5.L.154.244; 5.L.157.228; 5.L.157.229; 5.L.157.230; 5.L.157.231; 5.L.157.236;
5.L.157.237; 5.L.157.238; 5.L.157.239; 5.L.157.154; 5.L.157.157; 5.L.157.166;
5.L.157.169; 5.L.157.172; 5.L.157.175; 5.L.157.240; 5.L.157.244; 5.L.166.228;
20 5.L.166.229; 5.L.166.230; 5.L.166.231; 5.L.166.236; 5.L.166.237; 5.L.166.238;
5.L.166.239; 5.L.166.154; 5.L.166.157; 5.L.166.166; 5.L.166.169; 5.L.166.172;
5.L.166.175; 5.L.166.240; 5.L.166.244; 5.L.169.228; 5.L.169.229; 5.L.169.230;
5.L.169.231; 5.L.169.236; 5.L.169.237; 5.L.169.238; 5.L.169.239; 5.L.169.154;
5.L.169.157; 5.L.169.166; 5.L.169.169; 5.L.169.172; 5.L.169.175; 5.L.169.240;
25 5.L.169.244; 5.L.172.228; 5.L.172.229; 5.L.172.230; 5.L.172.231; 5.L.172.236;
5.L.172.237; 5.L.172.238; 5.L.172.239; 5.L.172.154; 5.L.172.157; 5.L.172.166;
5.L.172.169; 5.L.172.172; 5.L.172.175; 5.L.172.240; 5.L.172.244; 5.L.175.228;
5.L.175.229; 5.L.175.230; 5.L.175.231; 5.L.175.236; 5.L.175.237; 5.L.175.238;
5.L.175.239; 5.L.175.154; 5.L.175.157; 5.L.175.166; 5.L.175.169; 5.L.175.172;
30 5.L.175.175; 5.L.175.240; 5.L.175.244; 5.L.240.228; 5.L.240.229; 5.L.240.230;

5.L.240.231; 5.L.240.236; 5.L.240.237; 5.L.240.238; 5.L.240.239; 5.L.240.154;
 5.L.240.157; 5.L.240.166; 5.L.240.169; 5.L.240.172; 5.L.240.175; 5.L.240.240;
 5.L.240.244; 5.L.244.228; 5.L.244.229; 5.L.244.230; 5.L.244.231; 5.L.244.236;
 5.L.244.237; 5.L.244.238; 5.L.244.239; 5.L.244.154; 5.L.244.157; 5.L.244.166;
 5 5.L.244.169; 5.L.244.172; 5.L.244.175; 5.L.244.240; 5.L.244.244;

Prodrugs of 5.O

5.O.228.228; 5.O.228.229; 5.O.228.230; 5.O.228.231; 5.O.228.236;
 5.O.228.237; 5.O.228.238; 5.O.228.239; 5.O.228.154; 5.O.228.157;
 10 5.O.228.166; 5.O.228.169; 5.O.228.172; 5.O.228.175; 5.O.228.240;
 5.O.228.244; 5.O.229.228; 5.O.229.229; 5.O.229.230; 5.O.229.231;
 5.O.229.236; 5.O.229.237; 5.O.229.238; 5.O.229.239; 5.O.229.154;
 5.O.229.157; 5.O.229.166; 5.O.229.169; 5.O.229.172; 5.O.229.175;
 5.O.229.240; 5.O.229.244; 5.O.230.228; 5.O.230.229; 5.O.230.230;
 15 5.O.230.231; 5.O.230.236; 5.O.230.237; 5.O.230.238; 5.O.230.239;
 5.O.230.154; 5.O.230.157; 5.O.230.166; 5.O.230.169; 5.O.230.172;
 5.O.230.175; 5.O.230.240; 5.O.230.244; 5.O.231.228; 5.O.231.229;
 5.O.231.230; 5.O.231.231; 5.O.231.236; 5.O.231.237; 5.O.231.238;
 5.O.231.239; 5.O.231.154; 5.O.231.157; 5.O.231.166; 5.O.231.169;
 20 5.O.231.172; 5.O.231.175; 5.O.231.240; 5.O.231.244; 5.O.236.228;
 5.O.236.229; 5.O.236.230; 5.O.236.231; 5.O.236.236; 5.O.236.237;
 5.O.236.238; 5.O.236.239; 5.O.236.154; 5.O.236.157; 5.O.236.166;
 5.O.236.169; 5.O.236.172; 5.O.236.175; 5.O.236.240; 5.O.236.244;
 5.O.237.228; 5.O.237.229; 5.O.237.230; 5.O.237.231; 5.O.237.236;
 25 5.O.237.237; 5.O.237.238; 5.O.237.239; 5.O.237.154; 5.O.237.157;
 5.O.237.166; 5.O.237.169; 5.O.237.172; 5.O.237.175; 5.O.237.240;
 5.O.237.244; 5.O.238.228; 5.O.238.229; 5.O.238.230; 5.O.238.231;
 5.O.238.236; 5.O.238.237; 5.O.238.238; 5.O.238.239; 5.O.238.154;
 5.O.238.157; 5.O.238.166; 5.O.238.169; 5.O.238.172; 5.O.238.175;
 30 5.O.238.240; 5.O.238.244; 5.O.239.228; 5.O.239.229; 5.O.239.230;

5.O.239.231; 5.O.239.236; 5.O.239.237; 5.O.239.238; 5.O.239.239;
5.O.239.154; 5.O.239.157; 5.O.239.166; 5.O.239.169; 5.O.239.172;
5.O.239.175; 5.O.239.240; 5.O.239.244; 5.O.154.228; 5.O.154.229;
5.O.154.230; 5.O.154.231; 5.O.154.236; 5.O.154.237; 5.O.154.238;
5 5.O.154.239; 5.O.154.154; 5.O.154.157; 5.O.154.166; 5.O.154.169;
5.O.154.172; 5.O.154.175; 5.O.154.240; 5.O.154.244; 5.O.157.228;
5.O.157.229; 5.O.157.230; 5.O.157.231; 5.O.157.236; 5.O.157.237;
5.O.157.238; 5.O.157.239; 5.O.157.154; 5.O.157.157; 5.O.157.166;
5.O.157.169; 5.O.157.172; 5.O.157.175; 5.O.157.240; 5.O.157.244;
10 5.O.166.228; 5.O.166.229; 5.O.166.230; 5.O.166.231; 5.O.166.236;
5.O.166.237; 5.O.166.238; 5.O.166.239; 5.O.166.154; 5.O.166.157;
5.O.166.166; 5.O.166.169; 5.O.166.172; 5.O.166.175; 5.O.166.240;
5.O.166.244; 5.O.169.228; 5.O.169.229; 5.O.169.230; 5.O.169.231;
5.O.169.236; 5.O.169.237; 5.O.169.238; 5.O.169.239; 5.O.169.154;
15 5.O.169.157; 5.O.169.166; 5.O.169.169; 5.O.169.172; 5.O.169.175;
5.O.169.240; 5.O.169.244; 5.O.172.228; 5.O.172.229; 5.O.172.230;
5.O.172.231; 5.O.172.236; 5.O.172.237; 5.O.172.238; 5.O.172.239;
5.O.172.154; 5.O.172.157; 5.O.172.166; 5.O.172.169; 5.O.172.172;
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20 5.O.175.230; 5.O.175.231; 5.O.175.236; 5.O.175.237; 5.O.175.238;
5.O.175.239; 5.O.175.154; 5.O.175.157; 5.O.175.166; 5.O.175.169;
5.O.175.172; 5.O.175.175; 5.O.175.240; 5.O.175.244; 5.O.240.228;
5.O.240.229; 5.O.240.230; 5.O.240.231; 5.O.240.236; 5.O.240.237;
5.O.240.238; 5.O.240.239; 5.O.240.154; 5.O.240.157; 5.O.240.166;
25 5.O.240.169; 5.O.240.172; 5.O.240.175; 5.O.240.240; 5.O.240.244;
5.O.244.228; 5.O.244.229; 5.O.244.230; 5.O.244.231; 5.O.244.236;
5.O.244.237; 5.O.244.238; 5.O.244.239; 5.O.244.154; 5.O.244.157;
5.O.244.166; 5.O.244.169; 5.O.244.172; 5.O.244.175; 5.O.244.240;
5.O.244.244;

Prodrugs of 5.P

5.P.228.228; 5.P.228.229; 5.P.228.230; 5.P.228.231; 5.P.228.236;
5.P.228.237; 5.P.228.238; 5.P.228.239; 5.P.228.154; 5.P.228.157; 5.P.228.166;
5.P.228.169; 5.P.228.172; 5.P.228.175; 5.P.228.240; 5.P.228.244; 5.P.229.228;
5 5.P.229.229; 5.P.229.230; 5.P.229.231; 5.P.229.236; 5.P.229.237; 5.P.229.238;
5.P.229.239; 5.P.229.154; 5.P.229.157; 5.P.229.166; 5.P.229.169; 5.P.229.172;
5.P.229.175; 5.P.229.240; 5.P.229.244; 5.P.230.228; 5.P.230.229; 5.P.230.230;
5.P.230.231; 5.P.230.236; 5.P.230.237; 5.P.230.238; 5.P.230.239; 5.P.230.154;
5.P.230.157; 5.P.230.166; 5.P.230.169; 5.P.230.172; 5.P.230.175; 5.P.230.240;
10 5.P.230.244; 5.P.231.228; 5.P.231.229; 5.P.231.230; 5.P.231.231; 5.P.231.236;
5.P.231.237; 5.P.231.238; 5.P.231.239; 5.P.231.154; 5.P.231.157; 5.P.231.166;
5.P.231.169; 5.P.231.172; 5.P.231.175; 5.P.231.240; 5.P.231.244; 5.P.236.228;
5.P.236.229; 5.P.236.230; 5.P.236.231; 5.P.236.236; 5.P.236.237; 5.P.236.238;
5.P.236.239; 5.P.236.154; 5.P.236.157; 5.P.236.166; 5.P.236.169; 5.P.236.172;
15 5.P.236.175; 5.P.236.240; 5.P.236.244; 5.P.237.228; 5.P.237.229; 5.P.237.230;
5.P.237.231; 5.P.237.236; 5.P.237.237; 5.P.237.238; 5.P.237.239; 5.P.237.154;
5.P.237.157; 5.P.237.166; 5.P.237.169; 5.P.237.172; 5.P.237.175; 5.P.237.240;
5.P.237.244; 5.P.238.228; 5.P.238.229; 5.P.238.230; 5.P.238.231; 5.P.238.236;
5.P.238.237; 5.P.238.238; 5.P.238.239; 5.P.238.154; 5.P.238.157; 5.P.238.166;
20 5.P.238.169; 5.P.238.172; 5.P.238.175; 5.P.238.240; 5.P.238.244; 5.P.239.228;
5.P.239.229; 5.P.239.230; 5.P.239.231; 5.P.239.236; 5.P.239.237; 5.P.239.238;
5.P.239.239; 5.P.239.154; 5.P.239.157; 5.P.239.166; 5.P.239.169; 5.P.239.172;
5.P.239.175; 5.P.239.240; 5.P.239.244; 5.P.154.228; 5.P.154.229; 5.P.154.230;
5.P.154.231; 5.P.154.236; 5.P.154.237; 5.P.154.238; 5.P.154.239; 5.P.154.154;
25 5.P.154.157; 5.P.154.166; 5.P.154.169; 5.P.154.172; 5.P.154.175; 5.P.154.240;
5.P.154.244; 5.P.157.228; 5.P.157.229; 5.P.157.230; 5.P.157.231; 5.P.157.236;
5.P.157.237; 5.P.157.238; 5.P.157.239; 5.P.157.154; 5.P.157.157; 5.P.157.166;
5.P.157.169; 5.P.157.172; 5.P.157.175; 5.P.157.240; 5.P.157.244; 5.P.166.228;
5.P.166.229; 5.P.166.230; 5.P.166.231; 5.P.166.236; 5.P.166.237; 5.P.166.238;
30 5.P.166.239; 5.P.166.154; 5.P.166.157; 5.P.166.166; 5.P.166.169; 5.P.166.172;

5.P.166.175; 5.P.166.240; 5.P.166.244; 5.P.169.228; 5.P.169.229; 5.P.169.230;
5.P.169.231; 5.P.169.236; 5.P.169.237; 5.P.169.238; 5.P.169.239; 5.P.169.154;
5.P.169.157; 5.P.169.166; 5.P.169.169; 5.P.169.172; 5.P.169.175; 5.P.169.240;
5.P.169.244; 5.P.172.228; 5.P.172.229; 5.P.172.230; 5.P.172.231; 5.P.172.236;
5 5.P.172.237; 5.P.172.238; 5.P.172.239; 5.P.172.154; 5.P.172.157; 5.P.172.166;
5.P.172.169; 5.P.172.172; 5.P.172.175; 5.P.172.240; 5.P.172.244; 5.P.175.228;
5.P.175.229; 5.P.175.230; 5.P.175.231; 5.P.175.236; 5.P.175.237; 5.P.175.238;
5.P.175.239; 5.P.175.154; 5.P.175.157; 5.P.175.166; 5.P.175.169; 5.P.175.172;
5.P.175.175; 5.P.175.240; 5.P.175.244; 5.P.240.228; 5.P.240.229; 5.P.240.230;
10 5.P.240.231; 5.P.240.236; 5.P.240.237; 5.P.240.238; 5.P.240.239; 5.P.240.154;
5.P.240.157; 5.P.240.166; 5.P.240.169; 5.P.240.172; 5.P.240.175; 5.P.240.240;
5.P.240.244; 5.P.244.228; 5.P.244.229; 5.P.244.230; 5.P.244.231; 5.P.244.236;
5.P.244.237; 5.P.244.238; 5.P.244.239; 5.P.244.154; 5.P.244.157; 5.P.244.166;
5.P.244.169; 5.P.244.172; 5.P.244.175; 5.P.244.240; 5.P.244.244;

15

Prodrugs of 5.U

5.U.228.228; 5.U.228.229; 5.U.228.230; 5.U.228.231; 5.U.228.236;
5.U.228.237; 5.U.228.238; 5.U.228.239; 5.U.228.154; 5.U.228.157;
5.U.228.166; 5.U.228.169; 5.U.228.172; 5.U.228.175; 5.U.228.240;
20 5.U.228.244; 5.U.229.228; 5.U.229.229; 5.U.229.230; 5.U.229.231;
5.U.229.236; 5.U.229.237; 5.U.229.238; 5.U.229.239; 5.U.229.154;
5.U.229.157; 5.U.229.166; 5.U.229.169; 5.U.229.172; 5.U.229.175;
5.U.229.240; 5.U.229.244; 5.U.230.228; 5.U.230.229; 5.U.230.230;
5.U.230.231; 5.U.230.236; 5.U.230.237; 5.U.230.238; 5.U.230.239;
25 5.U.230.154; 5.U.230.157; 5.U.230.166; 5.U.230.169; 5.U.230.172;
5.U.230.175; 5.U.230.240; 5.U.230.244; 5.U.231.228; 5.U.231.229;
5.U.231.230; 5.U.231.231; 5.U.231.236; 5.U.231.237; 5.U.231.238;
5.U.231.239; 5.U.231.154; 5.U.231.157; 5.U.231.166; 5.U.231.169;
5.U.231.172; 5.U.231.175; 5.U.231.240; 5.U.231.244; 5.U.236.228;
30 5.U.236.229; 5.U.236.230; 5.U.236.231; 5.U.236.236; 5.U.236.237;

5.U.236.238; 5.U.236.239; 5.U.236.154; 5.U.236.157; 5.U.236.166;
5.U.236.169; 5.U.236.172; 5.U.236.175; 5.U.236.240; 5.U.236.244;
5.U.237.228; 5.U.237.229; 5.U.237.230; 5.U.237.231; 5.U.237.236;
5.U.237.237; 5.U.237.238; 5.U.237.239; 5.U.237.154; 5.U.237.157;
5 5.U.237.166; 5.U.237.169; 5.U.237.172; 5.U.237.175; 5.U.237.240;
5.U.237.244; 5.U.238.228; 5.U.238.229; 5.U.238.230; 5.U.238.231;
5.U.238.236; 5.U.238.237; 5.U.238.238; 5.U.238.239; 5.U.238.154;
5.U.238.157; 5.U.238.166; 5.U.238.169; 5.U.238.172; 5.U.238.175;
5.U.238.240; 5.U.238.244; 5.U.239.228; 5.U.239.229; 5.U.239.230;
10 5.U.239.231; 5.U.239.236; 5.U.239.237; 5.U.239.238; 5.U.239.239;
5.U.239.154; 5.U.239.157; 5.U.239.166; 5.U.239.169; 5.U.239.172;
5.U.239.175; 5.U.239.240; 5.U.239.244; 5.U.154.228; 5.U.154.229;
5.U.154.230; 5.U.154.231; 5.U.154.236; 5.U.154.237; 5.U.154.238;
5.U.154.239; 5.U.154.154; 5.U.154.157; 5.U.154.166; 5.U.154.169;
15 5.U.154.172; 5.U.154.175; 5.U.154.240; 5.U.154.244; 5.U.157.228;
5.U.157.229; 5.U.157.230; 5.U.157.231; 5.U.157.236; 5.U.157.237;
5.U.157.238; 5.U.157.239; 5.U.157.154; 5.U.157.157; 5.U.157.166;
5.U.157.169; 5.U.157.172; 5.U.157.175; 5.U.157.240; 5.U.157.244;
5.U.166.228; 5.U.166.229; 5.U.166.230; 5.U.166.231; 5.U.166.236;
20 5.U.166.237; 5.U.166.238; 5.U.166.239; 5.U.166.154; 5.U.166.157;
5.U.166.166; 5.U.166.169; 5.U.166.172; 5.U.166.175; 5.U.166.240;
5.U.166.244; 5.U.169.228; 5.U.169.229; 5.U.169.230; 5.U.169.231;
5.U.169.236; 5.U.169.237; 5.U.169.238; 5.U.169.239; 5.U.169.154;
5.U.169.157; 5.U.169.166; 5.U.169.169; 5.U.169.172; 5.U.169.175;
25 5.U.169.240; 5.U.169.244; 5.U.172.228; 5.U.172.229; 5.U.172.230;
5.U.172.231; 5.U.172.236; 5.U.172.237; 5.U.172.238; 5.U.172.239;
5.U.172.154; 5.U.172.157; 5.U.172.166; 5.U.172.169; 5.U.172.172;
5.U.172.175; 5.U.172.240; 5.U.172.244; 5.U.175.228; 5.U.175.229;
5.U.175.230; 5.U.175.231; 5.U.175.236; 5.U.175.237; 5.U.175.238;
30 5.U.175.239; 5.U.175.154; 5.U.175.157; 5.U.175.166; 5.U.175.169;

5.U.175.172; 5.U.175.175; 5.U.175.240; 5.U.175.244; 5.U.240.228;
5.U.240.229; 5.U.240.230; 5.U.240.231; 5.U.240.236; 5.U.240.237;
5.U.240.238; 5.U.240.239; 5.U.240.154; 5.U.240.157; 5.U.240.166;
5.U.240.169; 5.U.240.172; 5.U.240.175; 5.U.240.240; 5.U.240.244;
5 5.U.244.228; 5.U.244.229; 5.U.244.230; 5.U.244.231; 5.U.244.236;
5.U.244.237; 5.U.244.238; 5.U.244.239; 5.U.244.154; 5.U.244.157;
5.U.244.166; 5.U.244.169; 5.U.244.172; 5.U.244.175; 5.U.244.240;
5.U.244.244;

10 Prodrugs of 5.W

5.W.228.228; 5.W.228.229; 5.W.228.230; 5.W.228.231; 5.W.228.236;
5.W.228.237; 5.W.228.238; 5.W.228.239; 5.W.228.154; 5.W.228.157;
5.W.228.166; 5.W.228.169; 5.W.228.172; 5.W.228.175; 5.W.228.240;
5.W.228.244; 5.W.229.228; 5.W.229.229; 5.W.229.230; 5.W.229.231;
15 5.W.229.236; 5.W.229.237; 5.W.229.238; 5.W.229.239; 5.W.229.154;
5.W.229.157; 5.W.229.166; 5.W.229.169; 5.W.229.172; 5.W.229.175;
5.W.229.240; 5.W.229.244; 5.W.230.228; 5.W.230.229; 5.W.230.230;
5.W.230.231; 5.W.230.236; 5.W.230.237; 5.W.230.238; 5.W.230.239;
5.W.230.154; 5.W.230.157; 5.W.230.166; 5.W.230.169; 5.W.230.172;
20 5.W.230.175; 5.W.230.240; 5.W.230.244; 5.W.231.228; 5.W.231.229;
5.W.231.230; 5.W.231.231; 5.W.231.236; 5.W.231.237; 5.W.231.238;
5.W.231.239; 5.W.231.154; 5.W.231.157; 5.W.231.166; 5.W.231.169;
5.W.231.172; 5.W.231.175; 5.W.231.240; 5.W.231.244; 5.W.236.228;
5.W.236.229; 5.W.236.230; 5.W.236.231; 5.W.236.236; 5.W.236.237;
25 5.W.236.238; 5.W.236.239; 5.W.236.154; 5.W.236.157; 5.W.236.166;
5.W.236.169; 5.W.236.172; 5.W.236.175; 5.W.236.240; 5.W.236.244;
5.W.237.228; 5.W.237.229; 5.W.237.230; 5.W.237.231; 5.W.237.236;
5.W.237.237; 5.W.237.238; 5.W.237.239; 5.W.237.154; 5.W.237.157;
5.W.237.166; 5.W.237.169; 5.W.237.172; 5.W.237.175; 5.W.237.240;
30 5.W.237.244; 5.W.238.228; 5.W.238.229; 5.W.238.230; 5.W.238.231;

5.W.238.236; 5.W.238.237; 5.W.238.238; 5.W.238.239; 5.W.238.154;
5.W.238.157; 5.W.238.166; 5.W.238.169; 5.W.238.172; 5.W.238.175;
5.W.238.240; 5.W.238.244; 5.W.239.228; 5.W.239.229; 5.W.239.230;
5.W.239.231; 5.W.239.236; 5.W.239.237; 5.W.239.238; 5.W.239.239;
5 5.W.239.154; 5.W.239.157; 5.W.239.166; 5.W.239.169; 5.W.239.172;
5.W.239.175; 5.W.239.240; 5.W.239.244; 5.W.154.228; 5.W.154.229;
5.W.154.230; 5.W.154.231; 5.W.154.236; 5.W.154.237; 5.W.154.238;
5.W.154.239; 5.W.154.154; 5.W.154.157; 5.W.154.166; 5.W.154.169;
5.W.154.172; 5.W.154.175; 5.W.154.240; 5.W.154.244; 5.W.157.228;
10 5.W.157.229; 5.W.157.230; 5.W.157.231; 5.W.157.236; 5.W.157.237;
5.W.157.238; 5.W.157.239; 5.W.157.154; 5.W.157.157; 5.W.157.166;
5.W.157.169; 5.W.157.172; 5.W.157.175; 5.W.157.240; 5.W.157.244;
5.W.166.228; 5.W.166.229; 5.W.166.230; 5.W.166.231; 5.W.166.236;
5.W.166.237; 5.W.166.238; 5.W.166.239; 5.W.166.154; 5.W.166.157;
15 5.W.166.166; 5.W.166.169; 5.W.166.172; 5.W.166.175; 5.W.166.240;
5.W.166.244; 5.W.169.228; 5.W.169.229; 5.W.169.230; 5.W.169.231;
5.W.169.236; 5.W.169.237; 5.W.169.238; 5.W.169.239; 5.W.169.154;
5.W.169.157; 5.W.169.166; 5.W.169.169; 5.W.169.172; 5.W.169.175;
5.W.169.240; 5.W.169.244; 5.W.172.228; 5.W.172.229; 5.W.172.230;
20 5.W.172.231; 5.W.172.236; 5.W.172.237; 5.W.172.238; 5.W.172.239;
5.W.172.154; 5.W.172.157; 5.W.172.166; 5.W.172.169; 5.W.172.172;
5.W.172.175; 5.W.172.240; 5.W.172.244; 5.W.175.228; 5.W.175.229;
5.W.175.230; 5.W.175.231; 5.W.175.236; 5.W.175.237; 5.W.175.238;
5.W.175.239; 5.W.175.154; 5.W.175.157; 5.W.175.166; 5.W.175.169;
25 5.W.175.172; 5.W.175.175; 5.W.175.240; 5.W.175.244; 5.W.240.228;
5.W.240.229; 5.W.240.230; 5.W.240.231; 5.W.240.236; 5.W.240.237;
5.W.240.238; 5.W.240.239; 5.W.240.154; 5.W.240.157; 5.W.240.166;
5.W.240.169; 5.W.240.172; 5.W.240.175; 5.W.240.240; 5.W.240.244;
5.W.244.228; 5.W.244.229; 5.W.244.230; 5.W.244.231; 5.W.244.236;
30 5.W.244.237; 5.W.244.238; 5.W.244.239; 5.W.244.154; 5.W.244.157;

5.W.244.166; 5.W.244.169; 5.W.244.172; 5.W.244.175; 5.W.244.240;
5.W.244.244;

Prodrugs of 5.Y

- 5 5.Y.228.228; 5.Y.228.229; 5.Y.228.230; 5.Y.228.231; 5.Y.228.236;
5.Y.228.237; 5.Y.228.238; 5.Y.228.239; 5.Y.228.154; 5.Y.228.157; 5.Y.228.166;
5.Y.228.169; 5.Y.228.172; 5.Y.228.175; 5.Y.228.240; 5.Y.228.244; 5.Y.229.228;
5.Y.229.229; 5.Y.229.230; 5.Y.229.231; 5.Y.229.236; 5.Y.229.237; 5.Y.229.238;
5.Y.229.239; 5.Y.229.154; 5.Y.229.157; 5.Y.229.166; 5.Y.229.169; 5.Y.229.172;
10 5.Y.229.175; 5.Y.229.240; 5.Y.229.244; 5.Y.230.228; 5.Y.230.229; 5.Y.230.230;
5.Y.230.231; 5.Y.230.236; 5.Y.230.237; 5.Y.230.238; 5.Y.230.239; 5.Y.230.154;
5.Y.230.157; 5.Y.230.166; 5.Y.230.169; 5.Y.230.172; 5.Y.230.175; 5.Y.230.240;
5.Y.230.244; 5.Y.231.228; 5.Y.231.229; 5.Y.231.230; 5.Y.231.231; 5.Y.231.236;
5.Y.231.237; 5.Y.231.238; 5.Y.231.239; 5.Y.231.154; 5.Y.231.157; 5.Y.231.166;
15 5.Y.231.169; 5.Y.231.172; 5.Y.231.175; 5.Y.231.240; 5.Y.231.244; 5.Y.236.228;
5.Y.236.229; 5.Y.236.230; 5.Y.236.231; 5.Y.236.236; 5.Y.236.237; 5.Y.236.238;
5.Y.236.239; 5.Y.236.154; 5.Y.236.157; 5.Y.236.166; 5.Y.236.169; 5.Y.236.172;
5.Y.236.175; 5.Y.236.240; 5.Y.236.244; 5.Y.237.228; 5.Y.237.229; 5.Y.237.230;
5.Y.237.231; 5.Y.237.236; 5.Y.237.237; 5.Y.237.238; 5.Y.237.239; 5.Y.237.154;
20 5.Y.237.157; 5.Y.237.166; 5.Y.237.169; 5.Y.237.172; 5.Y.237.175; 5.Y.237.240;
5.Y.237.244; 5.Y.238.228; 5.Y.238.229; 5.Y.238.230; 5.Y.238.231; 5.Y.238.236;
5.Y.238.237; 5.Y.238.238; 5.Y.238.239; 5.Y.238.154; 5.Y.238.157; 5.Y.238.166;
5.Y.238.169; 5.Y.238.172; 5.Y.238.175; 5.Y.238.240; 5.Y.238.244; 5.Y.239.228;
5.Y.239.229; 5.Y.239.230; 5.Y.239.231; 5.Y.239.236; 5.Y.239.237; 5.Y.239.238;
25 5.Y.239.239; 5.Y.239.154; 5.Y.239.157; 5.Y.239.166; 5.Y.239.169; 5.Y.239.172;
5.Y.239.175; 5.Y.239.240; 5.Y.239.244; 5.Y.154.228; 5.Y.154.229; 5.Y.154.230;
5.Y.154.231; 5.Y.154.236; 5.Y.154.237; 5.Y.154.238; 5.Y.154.239; 5.Y.154.154;
5.Y.154.157; 5.Y.154.166; 5.Y.154.169; 5.Y.154.172; 5.Y.154.175; 5.Y.154.240;
5.Y.154.244; 5.Y.157.228; 5.Y.157.229; 5.Y.157.230; 5.Y.157.231; 5.Y.157.236;
30 5.Y.157.237; 5.Y.157.238; 5.Y.157.239; 5.Y.157.154; 5.Y.157.157; 5.Y.157.166;

5.Y.157.169; 5.Y.157.172; 5.Y.157.175; 5.Y.157.240; 5.Y.157.244; 5.Y.166.228;
 5.Y.166.229; 5.Y.166.230; 5.Y.166.231; 5.Y.166.236; 5.Y.166.237; 5.Y.166.238;
 5.Y.166.239; 5.Y.166.154; 5.Y.166.157; 5.Y.166.166; 5.Y.166.169; 5.Y.166.172;
 5.Y.166.175; 5.Y.166.240; 5.Y.166.244; 5.Y.169.228; 5.Y.169.229; 5.Y.169.230;
 5 5.Y.169.231; 5.Y.169.236; 5.Y.169.237; 5.Y.169.238; 5.Y.169.239; 5.Y.169.154;
 5.Y.169.157; 5.Y.169.166; 5.Y.169.169; 5.Y.169.172; 5.Y.169.175; 5.Y.169.240;
 5.Y.169.244; 5.Y.172.228; 5.Y.172.229; 5.Y.172.230; 5.Y.172.231; 5.Y.172.236;
 5.Y.172.237; 5.Y.172.238; 5.Y.172.239; 5.Y.172.154; 5.Y.172.157; 5.Y.172.166;
 5.Y.172.169; 5.Y.172.172; 5.Y.172.175; 5.Y.172.240; 5.Y.172.244; 5.Y.175.228;
 10 5.Y.175.229; 5.Y.175.230; 5.Y.175.231; 5.Y.175.236; 5.Y.175.237; 5.Y.175.238;
 5.Y.175.239; 5.Y.175.154; 5.Y.175.157; 5.Y.175.166; 5.Y.175.169; 5.Y.175.172;
 5.Y.175.175; 5.Y.175.240; 5.Y.175.244; 5.Y.240.228; 5.Y.240.229; 5.Y.240.230;
 5.Y.240.231; 5.Y.240.236; 5.Y.240.237; 5.Y.240.238; 5.Y.240.239; 5.Y.240.154;
 5.Y.240.157; 5.Y.240.166; 5.Y.240.169; 5.Y.240.172; 5.Y.240.175; 5.Y.240.240;
 15 5.Y.240.244; 5.Y.244.228; 5.Y.244.229; 5.Y.244.230; 5.Y.244.231; 5.Y.244.236;
 5.Y.244.237; 5.Y.244.238; 5.Y.244.239; 5.Y.244.154; 5.Y.244.157; 5.Y.244.166;
 5.Y.244.169; 5.Y.244.172; 5.Y.244.175; 5.Y.244.240; 5.Y.244.244;

Prodrugs of 6.B

20 6.B.228.228; 6.B.228.229; 6.B.228.230; 6.B.228.231; 6.B.228.236;
 6.B.228.237; 6.B.228.238; 6.B.228.239; 6.B.228.154; 6.B.228.157; 6.B.228.166;
 6.B.228.169; 6.B.228.172; 6.B.228.175; 6.B.228.240; 6.B.228.244; 6.B.229.228;
 6.B.229.229; 6.B.229.230; 6.B.229.231; 6.B.229.236; 6.B.229.237; 6.B.229.238;
 6.B.229.239; 6.B.229.154; 6.B.229.157; 6.B.229.166; 6.B.229.169; 6.B.229.172;
 25 6.B.229.175; 6.B.229.240; 6.B.229.244; 6.B.230.228; 6.B.230.229; 6.B.230.230;
 6.B.230.231; 6.B.230.236; 6.B.230.237; 6.B.230.238; 6.B.230.239; 6.B.230.154;
 6.B.230.157; 6.B.230.166; 6.B.230.169; 6.B.230.172; 6.B.230.175; 6.B.230.240;
 6.B.230.244; 6.B.231.228; 6.B.231.229; 6.B.231.230; 6.B.231.231; 6.B.231.236;
 6.B.231.237; 6.B.231.238; 6.B.231.239; 6.B.231.154; 6.B.231.157; 6.B.231.166;
 30 6.B.231.169; 6.B.231.172; 6.B.231.175; 6.B.231.240; 6.B.231.244; 6.B.236.228;

6.B.236.229; 6.B.236.230; 6.B.236.231; 6.B.236.236; 6.B.236.237; 6.B.236.238;
6.B.236.239; 6.B.236.154; 6.B.236.157; 6.B.236.166; 6.B.236.169; 6.B.236.172;
6.B.236.175; 6.B.236.240; 6.B.236.244; 6.B.237.228; 6.B.237.229; 6.B.237.230;
6.B.237.231; 6.B.237.236; 6.B.237.237; 6.B.237.238; 6.B.237.239; 6.B.237.154;
5 6.B.237.157; 6.B.237.166; 6.B.237.169; 6.B.237.172; 6.B.237.175; 6.B.237.240;
6.B.237.244; 6.B.238.228; 6.B.238.229; 6.B.238.230; 6.B.238.231; 6.B.238.236;
6.B.238.237; 6.B.238.238; 6.B.238.239; 6.B.238.154; 6.B.238.157; 6.B.238.166;
6.B.238.169; 6.B.238.172; 6.B.238.175; 6.B.238.240; 6.B.238.244; 6.B.239.228;
6.B.239.229; 6.B.239.230; 6.B.239.231; 6.B.239.236; 6.B.239.237; 6.B.239.238;
10 6.B.239.239; 6.B.239.154; 6.B.239.157; 6.B.239.166; 6.B.239.169; 6.B.239.172;
6.B.239.175; 6.B.239.240; 6.B.239.244; 6.B.154.228; 6.B.154.229; 6.B.154.230;
6.B.154.231; 6.B.154.236; 6.B.154.237; 6.B.154.238; 6.B.154.239; 6.B.154.154;
6.B.154.157; 6.B.154.166; 6.B.154.169; 6.B.154.172; 6.B.154.175; 6.B.154.240;
6.B.154.244; 6.B.157.228; 6.B.157.229; 6.B.157.230; 6.B.157.231; 6.B.157.236;
15 6.B.157.237; 6.B.157.238; 6.B.157.239; 6.B.157.154; 6.B.157.157; 6.B.157.166;
6.B.157.169; 6.B.157.172; 6.B.157.175; 6.B.157.240; 6.B.157.244; 6.B.166.228;
6.B.166.229; 6.B.166.230; 6.B.166.231; 6.B.166.236; 6.B.166.237; 6.B.166.238;
6.B.166.239; 6.B.166.154; 6.B.166.157; 6.B.166.166; 6.B.166.169; 6.B.166.172;
6.B.166.175; 6.B.166.240; 6.B.166.244; 6.B.169.228; 6.B.169.229; 6.B.169.230;
20 6.B.169.231; 6.B.169.236; 6.B.169.237; 6.B.169.238; 6.B.169.239; 6.B.169.154;
6.B.169.157; 6.B.169.166; 6.B.169.169; 6.B.169.172; 6.B.169.175; 6.B.169.240;
6.B.169.244; 6.B.172.228; 6.B.172.229; 6.B.172.230; 6.B.172.231; 6.B.172.236;
6.B.172.237; 6.B.172.238; 6.B.172.239; 6.B.172.154; 6.B.172.157; 6.B.172.166;
6.B.172.169; 6.B.172.172; 6.B.172.175; 6.B.172.240; 6.B.172.244; 6.B.175.228;
25 6.B.175.229; 6.B.175.230; 6.B.175.231; 6.B.175.236; 6.B.175.237; 6.B.175.238;
6.B.175.239; 6.B.175.154; 6.B.175.157; 6.B.175.166; 6.B.175.169; 6.B.175.172;
6.B.175.175; 6.B.175.240; 6.B.175.244; 6.B.240.228; 6.B.240.229; 6.B.240.230;
6.B.240.231; 6.B.240.236; 6.B.240.237; 6.B.240.238; 6.B.240.239; 6.B.240.154;
6.B.240.157; 6.B.240.166; 6.B.240.169; 6.B.240.172; 6.B.240.175; 6.B.240.240;
30 6.B.240.244; 6.B.244.228; 6.B.244.229; 6.B.244.230; 6.B.244.231; 6.B.244.236;

6.B.244.237; 6.B.244.238; 6.B.244.239; 6.B.244.154; 6.B.244.157; 6.B.244.166;
6.B.244.169; 6.B.244.172; 6.B.244.175; 6.B.244.240; 6.B.244.244;

Prodrugs of 6.D

- 5 6.D.228.228; 6.D.228.229; 6.D.228.230; 6.D.228.231; 6.D.228.236;
6.D.228.237; 6.D.228.238; 6.D.228.239; 6.D.228.154; 6.D.228.157;
6.D.228.166; 6.D.228.169; 6.D.228.172; 6.D.228.175; 6.D.228.240;
6.D.228.244; 6.D.229.228; 6.D.229.229; 6.D.229.230; 6.D.229.231;
6.D.229.236; 6.D.229.237; 6.D.229.238; 6.D.229.239; 6.D.229.154;
10 6.D.229.157; 6.D.229.166; 6.D.229.169; 6.D.229.172; 6.D.229.175;
6.D.229.240; 6.D.229.244; 6.D.230.228; 6.D.230.229; 6.D.230.230;
6.D.230.231; 6.D.230.236; 6.D.230.237; 6.D.230.238; 6.D.230.239;
6.D.230.154; 6.D.230.157; 6.D.230.166; 6.D.230.169; 6.D.230.172;
6.D.230.175; 6.D.230.240; 6.D.230.244; 6.D.231.228; 6.D.231.229;
15 6.D.231.230; 6.D.231.231; 6.D.231.236; 6.D.231.237; 6.D.231.238;
6.D.231.239; 6.D.231.154; 6.D.231.157; 6.D.231.166; 6.D.231.169;
6.D.231.172; 6.D.231.175; 6.D.231.240; 6.D.231.244; 6.D.236.228;
6.D.236.229; 6.D.236.230; 6.D.236.231; 6.D.236.236; 6.D.236.237;
6.D.236.238; 6.D.236.239; 6.D.236.154; 6.D.236.157; 6.D.236.166;
20 6.D.236.169; 6.D.236.172; 6.D.236.175; 6.D.236.240; 6.D.236.244;
6.D.237.228; 6.D.237.229; 6.D.237.230; 6.D.237.231; 6.D.237.236;
6.D.237.237; 6.D.237.238; 6.D.237.239; 6.D.237.154; 6.D.237.157;
6.D.237.166; 6.D.237.169; 6.D.237.172; 6.D.237.175; 6.D.237.240;
6.D.237.244; 6.D.238.228; 6.D.238.229; 6.D.238.230; 6.D.238.231;
25 6.D.238.236; 6.D.238.237; 6.D.238.238; 6.D.238.239; 6.D.238.154;
6.D.238.157; 6.D.238.166; 6.D.238.169; 6.D.238.172; 6.D.238.175;
6.D.238.240; 6.D.238.244; 6.D.239.228; 6.D.239.229; 6.D.239.230;
6.D.239.231; 6.D.239.236; 6.D.239.237; 6.D.239.238; 6.D.239.239;
6.D.239.154; 6.D.239.157; 6.D.239.166; 6.D.239.169; 6.D.239.172;
30 6.D.239.175; 6.D.239.240; 6.D.239.244; 6.D.154.228; 6.D.154.229;

- 6.D.154.230; 6.D.154.231; 6.D.154.236; 6.D.154.237; 6.D.154.238;
6.D.154.239; 6.D.154.154; 6.D.154.157; 6.D.154.166; 6.D.154.169;
6.D.154.172; 6.D.154.175; 6.D.154.240; 6.D.154.244; 6.D.157.228;
6.D.157.229; 6.D.157.230; 6.D.157.231; 6.D.157.236; 6.D.157.237;
5 6.D.157.238; 6.D.157.239; 6.D.157.154; 6.D.157.157; 6.D.157.166;
6.D.157.169; 6.D.157.172; 6.D.157.175; 6.D.157.240; 6.D.157.244;
6.D.166.228; 6.D.166.229; 6.D.166.230; 6.D.166.231; 6.D.166.236;
6.D.166.237; 6.D.166.238; 6.D.166.239; 6.D.166.154; 6.D.166.157;
6.D.166.166; 6.D.166.169; 6.D.166.172; 6.D.166.175; 6.D.166.240;
10 6.D.166.244; 6.D.169.228; 6.D.169.229; 6.D.169.230; 6.D.169.231;
6.D.169.236; 6.D.169.237; 6.D.169.238; 6.D.169.239; 6.D.169.154;
6.D.169.157; 6.D.169.166; 6.D.169.169; 6.D.169.172; 6.D.169.175;
6.D.169.240; 6.D.169.244; 6.D.172.228; 6.D.172.229; 6.D.172.230;
6.D.172.231; 6.D.172.236; 6.D.172.237; 6.D.172.238; 6.D.172.239;
15 6.D.172.154; 6.D.172.157; 6.D.172.166; 6.D.172.169; 6.D.172.172;
6.D.172.175; 6.D.172.240; 6.D.172.244; 6.D.175.228; 6.D.175.229;
6.D.175.230; 6.D.175.231; 6.D.175.236; 6.D.175.237; 6.D.175.238;
6.D.175.239; 6.D.175.154; 6.D.175.157; 6.D.175.166; 6.D.175.169;
6.D.175.172; 6.D.175.175; 6.D.175.240; 6.D.175.244; 6.D.240.228;
20 6.D.240.229; 6.D.240.230; 6.D.240.231; 6.D.240.236; 6.D.240.237;
6.D.240.238; 6.D.240.239; 6.D.240.154; 6.D.240.157; 6.D.240.166;
6.D.240.169; 6.D.240.172; 6.D.240.175; 6.D.240.240; 6.D.240.244;
6.D.244.228; 6.D.244.229; 6.D.244.230; 6.D.244.231; 6.D.244.236;
6.D.244.237; 6.D.244.238; 6.D.244.239; 6.D.244.154; 6.D.244.157;
25 6.D.244.166; 6.D.244.169; 6.D.244.172; 6.D.244.175; 6.D.244.240;
6.D.244.244;

Prodrugs of 6.E

- 6.E.228.228; 6.E.228.229; 6.E.228.230; 6.E.228.231; 6.E.228.236;
30 6.E.228.237; 6.E.228.238; 6.E.228.239; 6.E.228.154; 6.E.228.157; 6.E.228.166;

6.E.228.169; 6.E.228.172; 6.E.228.175; 6.E.228.240; 6.E.228.244; 6.E.229.228;
6.E.229.229; 6.E.229.230; 6.E.229.231; 6.E.229.236; 6.E.229.237; 6.E.229.238;
6.E.229.239; 6.E.229.154; 6.E.229.157; 6.E.229.166; 6.E.229.169; 6.E.229.172;
6.E.229.175; 6.E.229.240; 6.E.229.244; 6.E.230.228; 6.E.230.229; 6.E.230.230;
5 6.E.230.231; 6.E.230.236; 6.E.230.237; 6.E.230.238; 6.E.230.239; 6.E.230.154;
6.E.230.157; 6.E.230.166; 6.E.230.169; 6.E.230.172; 6.E.230.175; 6.E.230.240;
6.E.230.244; 6.E.231.228; 6.E.231.229; 6.E.231.230; 6.E.231.231; 6.E.231.236;
6.E.231.237; 6.E.231.238; 6.E.231.239; 6.E.231.154; 6.E.231.157; 6.E.231.166;
6.E.231.169; 6.E.231.172; 6.E.231.175; 6.E.231.240; 6.E.231.244; 6.E.236.228;
10 6.E.236.229; 6.E.236.230; 6.E.236.231; 6.E.236.236; 6.E.236.237; 6.E.236.238;
6.E.236.239; 6.E.236.154; 6.E.236.157; 6.E.236.166; 6.E.236.169; 6.E.236.172;
6.E.236.175; 6.E.236.240; 6.E.236.244; 6.E.237.228; 6.E.237.229; 6.E.237.230;
6.E.237.231; 6.E.237.236; 6.E.237.237; 6.E.237.238; 6.E.237.239; 6.E.237.154;
6.E.237.157; 6.E.237.166; 6.E.237.169; 6.E.237.172; 6.E.237.175; 6.E.237.240;
15 6.E.237.244; 6.E.238.228; 6.E.238.229; 6.E.238.230; 6.E.238.231; 6.E.238.236;
6.E.238.237; 6.E.238.238; 6.E.238.239; 6.E.238.154; 6.E.238.157; 6.E.238.166;
6.E.238.169; 6.E.238.172; 6.E.238.175; 6.E.238.240; 6.E.238.244; 6.E.239.228;
6.E.239.229; 6.E.239.230; 6.E.239.231; 6.E.239.236; 6.E.239.237; 6.E.239.238;
6.E.239.239; 6.E.239.154; 6.E.239.157; 6.E.239.166; 6.E.239.169; 6.E.239.172;
20 6.E.239.175; 6.E.239.240; 6.E.239.244; 6.E.154.228; 6.E.154.229; 6.E.154.230;
6.E.154.231; 6.E.154.236; 6.E.154.237; 6.E.154.238; 6.E.154.239; 6.E.154.154;
6.E.154.157; 6.E.154.166; 6.E.154.169; 6.E.154.172; 6.E.154.175; 6.E.154.240;
6.E.154.244; 6.E.157.228; 6.E.157.229; 6.E.157.230; 6.E.157.231; 6.E.157.236;
6.E.157.237; 6.E.157.238; 6.E.157.239; 6.E.157.154; 6.E.157.157; 6.E.157.166;
25 6.E.157.169; 6.E.157.172; 6.E.157.175; 6.E.157.240; 6.E.157.244; 6.E.166.228;
6.E.166.229; 6.E.166.230; 6.E.166.231; 6.E.166.236; 6.E.166.237; 6.E.166.238;
6.E.166.239; 6.E.166.154; 6.E.166.157; 6.E.166.166; 6.E.166.169; 6.E.166.172;
6.E.166.175; 6.E.166.240; 6.E.166.244; 6.E.169.228; 6.E.169.229; 6.E.169.230;
6.E.169.231; 6.E.169.236; 6.E.169.237; 6.E.169.238; 6.E.169.239; 6.E.169.154;
30 6.E.169.157; 6.E.169.166; 6.E.169.169; 6.E.169.172; 6.E.169.175; 6.E.169.240;

6.E.169.244; 6.E.172.228; 6.E.172.229; 6.E.172.230; 6.E.172.231; 6.E.172.236;
6.E.172.237; 6.E.172.238; 6.E.172.239; 6.E.172.154; 6.E.172.157; 6.E.172.166;
6.E.172.169; 6.E.172.172; 6.E.172.175; 6.E.172.240; 6.E.172.244; 6.E.175.228;
6.E.175.229; 6.E.175.230; 6.E.175.231; 6.E.175.236; 6.E.175.237; 6.E.175.238;
5 6.E.175.239; 6.E.175.154; 6.E.175.157; 6.E.175.166; 6.E.175.169; 6.E.175.172;
6.E.175.175; 6.E.175.240; 6.E.175.244; 6.E.240.228; 6.E.240.229; 6.E.240.230;
6.E.240.231; 6.E.240.236; 6.E.240.237; 6.E.240.238; 6.E.240.239; 6.E.240.154;
6.E.240.157; 6.E.240.166; 6.E.240.169; 6.E.240.172; 6.E.240.175; 6.E.240.240;
6.E.240.244; 6.E.244.228; 6.E.244.229; 6.E.244.230; 6.E.244.231; 6.E.244.236;
10 6.E.244.237; 6.E.244.238; 6.E.244.239; 6.E.244.154; 6.E.244.157; 6.E.244.166;
6.E.244.169; 6.E.244.172; 6.E.244.175; 6.E.244.240; 6.E.244.244;

Prodrugs of 6.G

6.G.228.228; 6.G.228.229; 6.G.228.230; 6.G.228.231; 6.G.228.236;
15 6.G.228.237; 6.G.228.238; 6.G.228.239; 6.G.228.154; 6.G.228.157;
6.G.228.166; 6.G.228.169; 6.G.228.172; 6.G.228.175; 6.G.228.240;
6.G.228.244; 6.G.229.228; 6.G.229.229; 6.G.229.230; 6.G.229.231;
6.G.229.236; 6.G.229.237; 6.G.229.238; 6.G.229.239; 6.G.229.154;
6.G.229.157; 6.G.229.166; 6.G.229.169; 6.G.229.172; 6.G.229.175;
20 6.G.229.240; 6.G.229.244; 6.G.230.228; 6.G.230.229; 6.G.230.230;
6.G.230.231; 6.G.230.236; 6.G.230.237; 6.G.230.238; 6.G.230.239;
6.G.230.154; 6.G.230.157; 6.G.230.166; 6.G.230.169; 6.G.230.172;
6.G.230.175; 6.G.230.240; 6.G.230.244; 6.G.231.228; 6.G.231.229;
6.G.231.230; 6.G.231.231; 6.G.231.236; 6.G.231.237; 6.G.231.238;
25 6.G.231.239; 6.G.231.154; 6.G.231.157; 6.G.231.166; 6.G.231.169;
6.G.231.172; 6.G.231.175; 6.G.231.240; 6.G.231.244; 6.G.236.228;
6.G.236.229; 6.G.236.230; 6.G.236.231; 6.G.236.236; 6.G.236.237;
6.G.236.238; 6.G.236.239; 6.G.236.154; 6.G.236.157; 6.G.236.166;
6.G.236.169; 6.G.236.172; 6.G.236.175; 6.G.236.240; 6.G.236.244;
30 6.G.237.228; 6.G.237.229; 6.G.237.230; 6.G.237.231; 6.G.237.236;

6.G.237.237; 6.G.237.238; 6.G.237.239; 6.G.237.154; 6.G.237.157;
6.G.237.166; 6.G.237.169; 6.G.237.172; 6.G.237.175; 6.G.237.240;
6.G.237.244; 6.G.238.228; 6.G.238.229; 6.G.238.230; 6.G.238.231;
6.G.238.236; 6.G.238.237; 6.G.238.238; 6.G.238.239; 6.G.238.154;
5 6.G.238.157; 6.G.238.166; 6.G.238.169; 6.G.238.172; 6.G.238.175;
6.G.238.240; 6.G.238.244; 6.G.239.228; 6.G.239.229; 6.G.239.230;
6.G.239.231; 6.G.239.236; 6.G.239.237; 6.G.239.238; 6.G.239.239;
6.G.239.154; 6.G.239.157; 6.G.239.166; 6.G.239.169; 6.G.239.172;
6.G.239.175; 6.G.239.240; 6.G.239.244; 6.G.154.228; 6.G.154.229;
10 6.G.154.230; 6.G.154.231; 6.G.154.236; 6.G.154.237; 6.G.154.238;
6.G.154.239; 6.G.154.154; 6.G.154.157; 6.G.154.166; 6.G.154.169;
6.G.154.172; 6.G.154.175; 6.G.154.240; 6.G.154.244; 6.G.157.228;
6.G.157.229; 6.G.157.230; 6.G.157.231; 6.G.157.236; 6.G.157.237;
6.G.157.238; 6.G.157.239; 6.G.157.154; 6.G.157.157; 6.G.157.166;
15 6.G.157.169; 6.G.157.172; 6.G.157.175; 6.G.157.240; 6.G.157.244;
6.G.166.228; 6.G.166.229; 6.G.166.230; 6.G.166.231; 6.G.166.236;
6.G.166.237; 6.G.166.238; 6.G.166.239; 6.G.166.154; 6.G.166.157;
6.G.166.166; 6.G.166.169; 6.G.166.172; 6.G.166.175; 6.G.166.240;
6.G.166.244; 6.G.169.228; 6.G.169.229; 6.G.169.230; 6.G.169.231;
20 6.G.169.236; 6.G.169.237; 6.G.169.238; 6.G.169.239; 6.G.169.154;
6.G.169.157; 6.G.169.166; 6.G.169.169; 6.G.169.172; 6.G.169.175;
6.G.169.240; 6.G.169.244; 6.G.172.228; 6.G.172.229; 6.G.172.230;
6.G.172.231; 6.G.172.236; 6.G.172.237; 6.G.172.238; 6.G.172.239;
6.G.172.154; 6.G.172.157; 6.G.172.166; 6.G.172.169; 6.G.172.172;
25 6.G.172.175; 6.G.172.240; 6.G.172.244; 6.G.175.228; 6.G.175.229;
6.G.175.230; 6.G.175.231; 6.G.175.236; 6.G.175.237; 6.G.175.238;
6.G.175.239; 6.G.175.154; 6.G.175.157; 6.G.175.166; 6.G.175.169;
6.G.175.172; 6.G.175.175; 6.G.175.240; 6.G.175.244; 6.G.240.228;
6.G.240.229; 6.G.240.230; 6.G.240.231; 6.G.240.236; 6.G.240.237;
30 6.G.240.238; 6.G.240.239; 6.G.240.154; 6.G.240.157; 6.G.240.166;

6.G.240.169; 6.G.240.172; 6.G.240.175; 6.G.240.240; 6.G.240.244;
6.G.244.228; 6.G.244.229; 6.G.244.230; 6.G.244.231; 6.G.244.236;
6.G.244.237; 6.G.244.238; 6.G.244.239; 6.G.244.154; 6.G.244.157;
6.G.244.166; 6.G.244.169; 6.G.244.172; 6.G.244.175; 6.G.244.240;
5 6.G.244.244;

Prodrugs of 6.I

6.I.228.228; 6.I.228.229; 6.I.228.230; 6.I.228.231; 6.I.228.236; 6.I.228.237;
6.I.228.238; 6.I.228.239; 6.I.228.154; 6.I.228.157; 6.I.228.166; 6.I.228.169;
10 6.I.228.172; 6.I.228.175; 6.I.228.240; 6.I.228.244; 6.I.229.228; 6.I.229.229;
6.I.229.230; 6.I.229.231; 6.I.229.236; 6.I.229.237; 6.I.229.238; 6.I.229.239;
6.I.229.154; 6.I.229.157; 6.I.229.166; 6.I.229.169; 6.I.229.172; 6.I.229.175;
6.I.229.240; 6.I.229.244; 6.I.230.228; 6.I.230.229; 6.I.230.230; 6.I.230.231;
6.I.230.236; 6.I.230.237; 6.I.230.238; 6.I.230.239; 6.I.230.154; 6.I.230.157;
15 6.I.230.166; 6.I.230.169; 6.I.230.172; 6.I.230.175; 6.I.230.240; 6.I.230.244;
6.I.231.228; 6.I.231.229; 6.I.231.230; 6.I.231.231; 6.I.231.236; 6.I.231.237;
6.I.231.238; 6.I.231.239; 6.I.231.154; 6.I.231.157; 6.I.231.166; 6.I.231.169;
6.I.231.172; 6.I.231.175; 6.I.231.240; 6.I.231.244; 6.I.236.228; 6.I.236.229;
6.I.236.230; 6.I.236.231; 6.I.236.236; 6.I.236.237; 6.I.236.238; 6.I.236.239;
20 6.I.236.154; 6.I.236.157; 6.I.236.166; 6.I.236.169; 6.I.236.172; 6.I.236.175;
6.I.236.240; 6.I.236.244; 6.I.237.228; 6.I.237.229; 6.I.237.230; 6.I.237.231;
6.I.237.236; 6.I.237.237; 6.I.237.238; 6.I.237.239; 6.I.237.154; 6.I.237.157;
6.I.237.166; 6.I.237.169; 6.I.237.172; 6.I.237.175; 6.I.237.240; 6.I.237.244;
6.I.238.228; 6.I.238.229; 6.I.238.230; 6.I.238.231; 6.I.238.236; 6.I.238.237;
25 6.I.238.238; 6.I.238.239; 6.I.238.154; 6.I.238.157; 6.I.238.166; 6.I.238.169;
6.I.238.172; 6.I.238.175; 6.I.238.240; 6.I.238.244; 6.I.239.228; 6.I.239.229;
6.I.239.230; 6.I.239.231; 6.I.239.236; 6.I.239.237; 6.I.239.238; 6.I.239.239;
6.I.239.154; 6.I.239.157; 6.I.239.166; 6.I.239.169; 6.I.239.172; 6.I.239.175;
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30 6.I.154.236; 6.I.154.237; 6.I.154.238; 6.I.154.239; 6.I.154.154; 6.I.154.157;

6.I.154.166; 6.I.154.169; 6.I.154.172; 6.I.154.175; 6.I.154.240; 6.I.154.244;
6.I.157.228; 6.I.157.229; 6.I.157.230; 6.I.157.231; 6.I.157.236; 6.I.157.237;
6.I.157.238; 6.I.157.239; 6.I.157.154; 6.I.157.157; 6.I.157.166; 6.I.157.169;
6.I.157.172; 6.I.157.175; 6.I.157.240; 6.I.157.244; 6.I.166.228; 6.I.166.229;
5 6.I.166.230; 6.I.166.231; 6.I.166.236; 6.I.166.237; 6.I.166.238; 6.I.166.239;
6.I.166.154; 6.I.166.157; 6.I.166.166; 6.I.166.169; 6.I.166.172; 6.I.166.175;
6.I.166.240; 6.I.166.244; 6.I.169.228; 6.I.169.229; 6.I.169.230; 6.I.169.231;
6.I.169.236; 6.I.169.237; 6.I.169.238; 6.I.169.239; 6.I.169.154; 6.I.169.157;
6.I.169.166; 6.I.169.169; 6.I.169.172; 6.I.169.175; 6.I.169.240; 6.I.169.244;
10 6.I.172.228; 6.I.172.229; 6.I.172.230; 6.I.172.231; 6.I.172.236; 6.I.172.237;
6.I.172.238; 6.I.172.239; 6.I.172.154; 6.I.172.157; 6.I.172.166; 6.I.172.169;
6.I.172.172; 6.I.172.175; 6.I.172.240; 6.I.172.244; 6.I.175.228; 6.I.175.229;
6.I.175.230; 6.I.175.231; 6.I.175.236; 6.I.175.237; 6.I.175.238; 6.I.175.239;
6.I.175.154; 6.I.175.157; 6.I.175.166; 6.I.175.169; 6.I.175.172; 6.I.175.175;
15 6.I.175.240; 6.I.175.244; 6.I.240.228; 6.I.240.229; 6.I.240.230; 6.I.240.231;
6.I.240.236; 6.I.240.237; 6.I.240.238; 6.I.240.239; 6.I.240.154; 6.I.240.157;
6.I.240.166; 6.I.240.169; 6.I.240.172; 6.I.240.175; 6.I.240.240; 6.I.240.244;
6.I.244.228; 6.I.244.229; 6.I.244.230; 6.I.244.231; 6.I.244.236; 6.I.244.237;
6.I.244.238; 6.I.244.239; 6.I.244.154; 6.I.244.157; 6.I.244.166; 6.I.244.169;
20 6.I.244.172; 6.I.244.175; 6.I.244.240; 6.I.244.244;

Prodrugs of 6.I

6.J.228.228; 6.J.228.229; 6.J.228.230; 6.J.228.231; 6.J.228.236; 6.J.228.237;
6.J.228.238; 6.J.228.239; 6.J.228.154; 6.J.228.157; 6.J.228.166; 6.J.228.169;
25 6.J.228.172; 6.J.228.175; 6.J.228.240; 6.J.228.244; 6.J.229.228; 6.J.229.229;
6.J.229.230; 6.J.229.231; 6.J.229.236; 6.J.229.237; 6.J.229.238; 6.J.229.239;
6.J.229.154; 6.J.229.157; 6.J.229.166; 6.J.229.169; 6.J.229.172; 6.J.229.175;
6.J.229.240; 6.J.229.244; 6.J.230.228; 6.J.230.229; 6.J.230.230; 6.J.230.231;
6.J.230.236; 6.J.230.237; 6.J.230.238; 6.J.230.239; 6.J.230.154; 6.J.230.157;
30 6.J.230.166; 6.J.230.169; 6.J.230.172; 6.J.230.175; 6.J.230.240; 6.J.230.244;

6.J.231.228; 6.J.231.229; 6.J.231.230; 6.J.231.231; 6.J.231.236; 6.J.231.237;
6.J.231.238; 6.J.231.239; 6.J.231.154; 6.J.231.157; 6.J.231.166; 6.J.231.169;
6.J.231.172; 6.J.231.175; 6.J.231.240; 6.J.231.244; 6.J.236.228; 6.J.236.229;
6.J.236.230; 6.J.236.231; 6.J.236.236; 6.J.236.237; 6.J.236.238; 6.J.236.239;
5 6.J.236.154; 6.J.236.157; 6.J.236.166; 6.J.236.169; 6.J.236.172; 6.J.236.175;
6.J.236.240; 6.J.236.244; 6.J.237.228; 6.J.237.229; 6.J.237.230; 6.J.237.231;
6.J.237.236; 6.J.237.237; 6.J.237.238; 6.J.237.239; 6.J.237.154; 6.J.237.157;
6.J.237.166; 6.J.237.169; 6.J.237.172; 6.J.237.175; 6.J.237.240; 6.J.237.244;
6.J.238.228; 6.J.238.229; 6.J.238.230; 6.J.238.231; 6.J.238.236; 6.J.238.237;
10 6.J.238.238; 6.J.238.239; 6.J.238.154; 6.J.238.157; 6.J.238.166; 6.J.238.169;
6.J.238.172; 6.J.238.175; 6.J.238.240; 6.J.238.244; 6.J.239.228; 6.J.239.229;
6.J.239.230; 6.J.239.231; 6.J.239.236; 6.J.239.237; 6.J.239.238; 6.J.239.239;
6.J.239.154; 6.J.239.157; 6.J.239.166; 6.J.239.169; 6.J.239.172; 6.J.239.175;
6.J.239.240; 6.J.239.244; 6.J.154.228; 6.J.154.229; 6.J.154.230; 6.J.154.231;
15 6.J.154.236; 6.J.154.237; 6.J.154.238; 6.J.154.239; 6.J.154.154; 6.J.154.157;
6.J.154.166; 6.J.154.169; 6.J.154.172; 6.J.154.175; 6.J.154.240; 6.J.154.244;
6.J.157.228; 6.J.157.229; 6.J.157.230; 6.J.157.231; 6.J.157.236; 6.J.157.237;
6.J.157.238; 6.J.157.239; 6.J.157.154; 6.J.157.157; 6.J.157.166; 6.J.157.169;
6.J.157.172; 6.J.157.175; 6.J.157.240; 6.J.157.244; 6.J.166.228; 6.J.166.229;
20 6.J.166.230; 6.J.166.231; 6.J.166.236; 6.J.166.237; 6.J.166.238; 6.J.166.239;
6.J.166.154; 6.J.166.157; 6.J.166.166; 6.J.166.169; 6.J.166.172; 6.J.166.175;
6.J.166.240; 6.J.166.244; 6.J.169.228; 6.J.169.229; 6.J.169.230; 6.J.169.231;
6.J.169.236; 6.J.169.237; 6.J.169.238; 6.J.169.239; 6.J.169.154; 6.J.169.157;
6.J.169.166; 6.J.169.169; 6.J.169.172; 6.J.169.175; 6.J.169.240; 6.J.169.244;
25 6.J.172.228; 6.J.172.229; 6.J.172.230; 6.J.172.231; 6.J.172.236; 6.J.172.237;
6.J.172.238; 6.J.172.239; 6.J.172.154; 6.J.172.157; 6.J.172.166; 6.J.172.169;
6.J.172.172; 6.J.172.175; 6.J.172.240; 6.J.172.244; 6.J.175.228; 6.J.175.229;
6.J.175.230; 6.J.175.231; 6.J.175.236; 6.J.175.237; 6.J.175.238; 6.J.175.239;
6.J.175.154; 6.J.175.157; 6.J.175.166; 6.J.175.169; 6.J.175.172; 6.J.175.175;
30 6.J.175.240; 6.J.175.244; 6.J.240.228; 6.J.240.229; 6.J.240.230; 6.J.240.231;

6.J.240.236; 6.J.240.237; 6.J.240.238; 6.J.240.239; 6.J.240.154; 6.J.240.157;
6.J.240.166; 6.J.240.169; 6.J.240.172; 6.J.240.175; 6.J.240.240; 6.J.240.244;
6.J.244.228; 6.J.244.229; 6.J.244.230; 6.J.244.231; 6.J.244.236; 6.J.244.237;
6.J.244.238; 6.J.244.239; 6.J.244.154; 6.J.244.157; 6.J.244.166; 6.J.244.169;
5 6.J.244.172; 6.J.244.175; 6.J.244.240; 6.J.244.244;

Prodrugs of 6.L

6.L.228.228; 6.L.228.229; 6.L.228.230; 6.L.228.231; 6.L.228.236;
6.L.228.237; 6.L.228.238; 6.L.228.239; 6.L.228.154; 6.L.228.157; 6.L.228.166;
10 6.L.228.169; 6.L.228.172; 6.L.228.175; 6.L.228.240; 6.L.228.244; 6.L.229.228;
6.L.229.229; 6.L.229.230; 6.L.229.231; 6.L.229.236; 6.L.229.237; 6.L.229.238;
6.L.229.239; 6.L.229.154; 6.L.229.157; 6.L.229.166; 6.L.229.169; 6.L.229.172;
6.L.229.175; 6.L.229.240; 6.L.229.244; 6.L.230.228; 6.L.230.229; 6.L.230.230;
6.L.230.231; 6.L.230.236; 6.L.230.237; 6.L.230.238; 6.L.230.239; 6.L.230.154;
15 6.L.230.157; 6.L.230.166; 6.L.230.169; 6.L.230.172; 6.L.230.175; 6.L.230.240;
6.L.230.244; 6.L.231.228; 6.L.231.229; 6.L.231.230; 6.L.231.231; 6.L.231.236;
6.L.231.237; 6.L.231.238; 6.L.231.239; 6.L.231.154; 6.L.231.157; 6.L.231.166;
6.L.231.169; 6.L.231.172; 6.L.231.175; 6.L.231.240; 6.L.231.244; 6.L.236.228;
6.L.236.229; 6.L.236.230; 6.L.236.231; 6.L.236.236; 6.L.236.237; 6.L.236.238;
20 6.L.236.239; 6.L.236.154; 6.L.236.157; 6.L.236.166; 6.L.236.169; 6.L.236.172;
6.L.236.175; 6.L.236.240; 6.L.236.244; 6.L.237.228; 6.L.237.229; 6.L.237.230;
6.L.237.231; 6.L.237.236; 6.L.237.237; 6.L.237.238; 6.L.237.239; 6.L.237.154;
6.L.237.157; 6.L.237.166; 6.L.237.169; 6.L.237.172; 6.L.237.175; 6.L.237.240;
6.L.237.244; 6.L.238.228; 6.L.238.229; 6.L.238.230; 6.L.238.231; 6.L.238.236;
25 6.L.238.237; 6.L.238.238; 6.L.238.239; 6.L.238.154; 6.L.238.157; 6.L.238.166;
6.L.238.169; 6.L.238.172; 6.L.238.175; 6.L.238.240; 6.L.238.244; 6.L.239.228;
6.L.239.229; 6.L.239.230; 6.L.239.231; 6.L.239.236; 6.L.239.237; 6.L.239.238;
6.L.239.239; 6.L.239.154; 6.L.239.157; 6.L.239.166; 6.L.239.169; 6.L.239.172;
6.L.239.175; 6.L.239.240; 6.L.239.244; 6.L.154.228; 6.L.154.229; 6.L.154.230;
30 6.L.154.231; 6.L.154.236; 6.L.154.237; 6.L.154.238; 6.L.154.239; 6.L.154.154;

- 6.L.154.157; 6.L.154.166; 6.L.154.169; 6.L.154.172; 6.L.154.175; 6.L.154.240;
6.L.154.244; 6.L.157.228; 6.L.157.229; 6.L.157.230; 6.L.157.231; 6.L.157.236;
6.L.157.237; 6.L.157.238; 6.L.157.239; 6.L.157.154; 6.L.157.157; 6.L.157.166;
6.L.157.169; 6.L.157.172; 6.L.157.175; 6.L.157.240; 6.L.157.244; 6.L.166.228;
5 6.L.166.229; 6.L.166.230; 6.L.166.231; 6.L.166.236; 6.L.166.237; 6.L.166.238;
6.L.166.239; 6.L.166.154; 6.L.166.157; 6.L.166.166; 6.L.166.169; 6.L.166.172;
6.L.166.175; 6.L.166.240; 6.L.166.244; 6.L.169.228; 6.L.169.229; 6.L.169.230;
6.L.169.231; 6.L.169.236; 6.L.169.237; 6.L.169.238; 6.L.169.239; 6.L.169.154;
6.L.169.157; 6.L.169.166; 6.L.169.169; 6.L.169.172; 6.L.169.175; 6.L.169.240;
10 6.L.169.244; 6.L.172.228; 6.L.172.229; 6.L.172.230; 6.L.172.231; 6.L.172.236;
6.L.172.237; 6.L.172.238; 6.L.172.239; 6.L.172.154; 6.L.172.157; 6.L.172.166;
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6.L.175.239; 6.L.175.154; 6.L.175.157; 6.L.175.166; 6.L.175.169; 6.L.175.172;
15 6.L.175.175; 6.L.175.240; 6.L.175.244; 6.L.240.228; 6.L.240.229; 6.L.240.230;
6.L.240.231; 6.L.240.236; 6.L.240.237; 6.L.240.238; 6.L.240.239; 6.L.240.154;
6.L.240.157; 6.L.240.166; 6.L.240.169; 6.L.240.172; 6.L.240.175; 6.L.240.240;
6.L.240.244; 6.L.244.228; 6.L.244.229; 6.L.244.230; 6.L.244.231; 6.L.244.236;
6.L.244.237; 6.L.244.238; 6.L.244.239; 6.L.244.154; 6.L.244.157; 6.L.244.166;
20 6.L.244.169; 6.L.244.172; 6.L.244.175; 6.L.244.240; 6.L.244.244;

Prodrugs of 6.O

- 6.O.228.228; 6.O.228.229; 6.O.228.230; 6.O.228.231; 6.O.228.236;
6.O.228.237; 6.O.228.238; 6.O.228.239; 6.O.228.154; 6.O.228.157;
25 6.O.228.166; 6.O.228.169; 6.O.228.172; 6.O.228.175; 6.O.228.240;
6.O.228.244; 6.O.229.228; 6.O.229.229; 6.O.229.230; 6.O.229.231;
6.O.229.236; 6.O.229.237; 6.O.229.238; 6.O.229.239; 6.O.229.154;
6.O.229.157; 6.O.229.166; 6.O.229.169; 6.O.229.172; 6.O.229.175;
6.O.229.240; 6.O.229.244; 6.O.230.228; 6.O.230.229; 6.O.230.230;
30 6.O.230.231; 6.O.230.236; 6.O.230.237; 6.O.230.238; 6.O.230.239;

6.O.230.154; 6.O.230.157; 6.O.230.166; 6.O.230.169; 6.O.230.172;
6.O.230.175; 6.O.230.240; 6.O.230.244; 6.O.231.228; 6.O.231.229;
6.O.231.230; 6.O.231.231; 6.O.231.236; 6.O.231.237; 6.O.231.238;
6.O.231.239; 6.O.231.154; 6.O.231.157; 6.O.231.166; 6.O.231.169;
5 6.O.231.172; 6.O.231.175; 6.O.231.240; 6.O.231.244; 6.O.236.228;
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6.O.236.238; 6.O.236.239; 6.O.236.154; 6.O.236.157; 6.O.236.166;
6.O.236.169; 6.O.236.172; 6.O.236.175; 6.O.236.240; 6.O.236.244;
6.O.237.228; 6.O.237.229; 6.O.237.230; 6.O.237.231; 6.O.237.236;
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6.O.238.157; 6.O.238.166; 6.O.238.169; 6.O.238.172; 6.O.238.175;
15 6.O.238.240; 6.O.238.244; 6.O.239.228; 6.O.239.229; 6.O.239.230;
6.O.239.231; 6.O.239.236; 6.O.239.237; 6.O.239.238; 6.O.239.239;
6.O.239.154; 6.O.239.157; 6.O.239.166; 6.O.239.169; 6.O.239.172;
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6.O.154.230; 6.O.154.231; 6.O.154.236; 6.O.154.237; 6.O.154.238;
20 6.O.154.239; 6.O.154.154; 6.O.154.157; 6.O.154.166; 6.O.154.169;
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6.O.157.238; 6.O.157.239; 6.O.157.154; 6.O.157.157; 6.O.157.166;
6.O.157.169; 6.O.157.172; 6.O.157.175; 6.O.157.240; 6.O.157.244;
25 6.O.166.228; 6.O.166.229; 6.O.166.230; 6.O.166.231; 6.O.166.236;
6.O.166.237; 6.O.166.238; 6.O.166.239; 6.O.166.154; 6.O.166.157;
6.O.166.166; 6.O.166.169; 6.O.166.172; 6.O.166.175; 6.O.166.240;
6.O.166.244; 6.O.169.228; 6.O.169.229; 6.O.169.230; 6.O.169.231;
6.O.169.236; 6.O.169.237; 6.O.169.238; 6.O.169.239; 6.O.169.154;
30 6.O.169.157; 6.O.169.166; 6.O.169.169; 6.O.169.172; 6.O.169.175;

6.O.169.240; 6.O.169.244; 6.O.172.228; 6.O.172.229; 6.O.172.230;
6.O.172.231; 6.O.172.236; 6.O.172.237; 6.O.172.238; 6.O.172.239;
6.O.172.154; 6.O.172.157; 6.O.172.166; 6.O.172.169; 6.O.172.172;
6.O.172.175; 6.O.172.240; 6.O.172.244; 6.O.175.228; 6.O.175.229;
5 6.O.175.230; 6.O.175.231; 6.O.175.236; 6.O.175.237; 6.O.175.238;
6.O.175.239; 6.O.175.154; 6.O.175.157; 6.O.175.166; 6.O.175.169;
6.O.175.172; 6.O.175.175; 6.O.175.240; 6.O.175.244; 6.O.240.228;
6.O.240.229; 6.O.240.230; 6.O.240.231; 6.O.240.236; 6.O.240.237;
6.O.240.238; 6.O.240.239; 6.O.240.154; 6.O.240.157; 6.O.240.166;
10 6.O.240.169; 6.O.240.172; 6.O.240.175; 6.O.240.240; 6.O.240.244;
6.O.244.228; 6.O.244.229; 6.O.244.230; 6.O.244.231; 6.O.244.236;
6.O.244.237; 6.O.244.238; 6.O.244.239; 6.O.244.154; 6.O.244.157;
6.O.244.166; 6.O.244.169; 6.O.244.172; 6.O.244.175; 6.O.244.240;
6.O.244.244;

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Prodrugs of 6.P

6.P.228.228; 6.P.228.229; 6.P.228.230; 6.P.228.231; 6.P.228.236;
6.P.228.237; 6.P.228.238; 6.P.228.239; 6.P.228.154; 6.P.228.157; 6.P.228.166;
6.P.228.169; 6.P.228.172; 6.P.228.175; 6.P.228.240; 6.P.228.244; 6.P.229.228;
20 6.P.229.229; 6.P.229.230; 6.P.229.231; 6.P.229.236; 6.P.229.237; 6.P.229.238;
6.P.229.239; 6.P.229.154; 6.P.229.157; 6.P.229.166; 6.P.229.169; 6.P.229.172;
6.P.229.175; 6.P.229.240; 6.P.229.244; 6.P.230.228; 6.P.230.229; 6.P.230.230;
6.P.230.231; 6.P.230.236; 6.P.230.237; 6.P.230.238; 6.P.230.239; 6.P.230.154;
6.P.230.157; 6.P.230.166; 6.P.230.169; 6.P.230.172; 6.P.230.175; 6.P.230.240;
25 6.P.230.244; 6.P.231.228; 6.P.231.229; 6.P.231.230; 6.P.231.231; 6.P.231.236;
6.P.231.237; 6.P.231.238; 6.P.231.239; 6.P.231.154; 6.P.231.157; 6.P.231.166;
6.P.231.169; 6.P.231.172; 6.P.231.175; 6.P.231.240; 6.P.231.244; 6.P.236.228;
6.P.236.229; 6.P.236.230; 6.P.236.231; 6.P.236.236; 6.P.236.237; 6.P.236.238;
6.P.236.239; 6.P.236.154; 6.P.236.157; 6.P.236.166; 6.P.236.169; 6.P.236.172;
30 6.P.236.175; 6.P.236.240; 6.P.236.244; 6.P.237.228; 6.P.237.229; 6.P.237.230;

6.P.237.231; 6.P.237.236; 6.P.237.237; 6.P.237.238; 6.P.237.239; 6.P.237.154;
6.P.237.157; 6.P.237.166; 6.P.237.169; 6.P.237.172; 6.P.237.175; 6.P.237.240;
6.P.237.244; 6.P.238.228; 6.P.238.229; 6.P.238.230; 6.P.238.231; 6.P.238.236;
6.P.238.237; 6.P.238.238; 6.P.238.239; 6.P.238.154; 6.P.238.157; 6.P.238.166;
5 6.P.238.169; 6.P.238.172; 6.P.238.175; 6.P.238.240; 6.P.238.244; 6.P.239.228;
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6.P.239.175; 6.P.239.240; 6.P.239.244; 6.P.154.228; 6.P.154.229; 6.P.154.230;
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10 6.P.154.157; 6.P.154.166; 6.P.154.169; 6.P.154.172; 6.P.154.175; 6.P.154.240;
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6.P.157.237; 6.P.157.238; 6.P.157.239; 6.P.157.154; 6.P.157.157; 6.P.157.166;
6.P.157.169; 6.P.157.172; 6.P.157.175; 6.P.157.240; 6.P.157.244; 6.P.166.228;
6.P.166.229; 6.P.166.230; 6.P.166.231; 6.P.166.236; 6.P.166.237; 6.P.166.238;
15 6.P.166.239; 6.P.166.154; 6.P.166.157; 6.P.166.166; 6.P.166.169; 6.P.166.172;
6.P.166.175; 6.P.166.240; 6.P.166.244; 6.P.169.228; 6.P.169.229; 6.P.169.230;
6.P.169.231; 6.P.169.236; 6.P.169.237; 6.P.169.238; 6.P.169.239; 6.P.169.154;
6.P.169.157; 6.P.169.166; 6.P.169.169; 6.P.169.172; 6.P.169.175; 6.P.169.240;
6.P.169.244; 6.P.172.228; 6.P.172.229; 6.P.172.230; 6.P.172.231; 6.P.172.236;
20 6.P.172.237; 6.P.172.238; 6.P.172.239; 6.P.172.154; 6.P.172.157; 6.P.172.166;
6.P.172.169; 6.P.172.172; 6.P.172.175; 6.P.172.240; 6.P.172.244; 6.P.175.228;
6.P.175.229; 6.P.175.230; 6.P.175.231; 6.P.175.236; 6.P.175.237; 6.P.175.238;
6.P.175.239; 6.P.175.154; 6.P.175.157; 6.P.175.166; 6.P.175.169; 6.P.175.172;
6.P.175.175; 6.P.175.240; 6.P.175.244; 6.P.240.228; 6.P.240.229; 6.P.240.230;
25 6.P.240.231; 6.P.240.236; 6.P.240.237; 6.P.240.238; 6.P.240.239; 6.P.240.154;
6.P.240.157; 6.P.240.166; 6.P.240.169; 6.P.240.172; 6.P.240.175; 6.P.240.240;
6.P.240.244; 6.P.244.228; 6.P.244.229; 6.P.244.230; 6.P.244.231; 6.P.244.236;
6.P.244.237; 6.P.244.238; 6.P.244.239; 6.P.244.154; 6.P.244.157; 6.P.244.166;
6.P.244.169; 6.P.244.172; 6.P.244.175; 6.P.244.240; 6.P.244.244;

Prodrugs of 6.U

- 6.U.228.228; 6.U.228.229; 6.U.228.230; 6.U.228.231; 6.U.228.236;
6.U.228.237; 6.U.228.238; 6.U.228.239; 6.U.228.154; 6.U.228.157;
6.U.228.166; 6.U.228.169; 6.U.228.172; 6.U.228.175; 6.U.228.240;
5 6.U.228.244; 6.U.229.228; 6.U.229.229; 6.U.229.230; 6.U.229.231;
6.U.229.236; 6.U.229.237; 6.U.229.238; 6.U.229.239; 6.U.229.154;
6.U.229.157; 6.U.229.166; 6.U.229.169; 6.U.229.172; 6.U.229.175;
6.U.229.240; 6.U.229.244; 6.U.230.228; 6.U.230.229; 6.U.230.230;
6.U.230.231; 6.U.230.236; 6.U.230.237; 6.U.230.238; 6.U.230.239;
10 6.U.230.154; 6.U.230.157; 6.U.230.166; 6.U.230.169; 6.U.230.172;
6.U.230.175; 6.U.230.240; 6.U.230.244; 6.U.231.228; 6.U.231.229;
6.U.231.230; 6.U.231.231; 6.U.231.236; 6.U.231.237; 6.U.231.238;
6.U.231.239; 6.U.231.154; 6.U.231.157; 6.U.231.166; 6.U.231.169;
6.U.231.172; 6.U.231.175; 6.U.231.240; 6.U.231.244; 6.U.236.228;
15 6.U.236.229; 6.U.236.230; 6.U.236.231; 6.U.236.236; 6.U.236.237;
6.U.236.238; 6.U.236.239; 6.U.236.154; 6.U.236.157; 6.U.236.166;
6.U.236.169; 6.U.236.172; 6.U.236.175; 6.U.236.240; 6.U.236.244;
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6.U.237.237; 6.U.237.238; 6.U.237.239; 6.U.237.154; 6.U.237.157;
20 6.U.237.166; 6.U.237.169; 6.U.237.172; 6.U.237.175; 6.U.237.240;
6.U.237.244; 6.U.238.228; 6.U.238.229; 6.U.238.230; 6.U.238.231;
6.U.238.236; 6.U.238.237; 6.U.238.238; 6.U.238.239; 6.U.238.154;
6.U.238.157; 6.U.238.166; 6.U.238.169; 6.U.238.172; 6.U.238.175;
6.U.238.240; 6.U.238.244; 6.U.239.228; 6.U.239.229; 6.U.239.230;
25 6.U.239.231; 6.U.239.236; 6.U.239.237; 6.U.239.238; 6.U.239.239;
6.U.239.154; 6.U.239.157; 6.U.239.166; 6.U.239.169; 6.U.239.172;
6.U.239.175; 6.U.239.240; 6.U.239.244; 6.U.154.228; 6.U.154.229;
6.U.154.230; 6.U.154.231; 6.U.154.236; 6.U.154.237; 6.U.154.238;
6.U.154.239; 6.U.154.154; 6.U.154.157; 6.U.154.166; 6.U.154.169;
30 6.U.154.172; 6.U.154.175; 6.U.154.240; 6.U.154.244; 6.U.157.228;

- 6.U.157.229; 6.U.157.230; 6.U.157.231; 6.U.157.236; 6.U.157.237;
6.U.157.238; 6.U.157.239; 6.U.157.154; 6.U.157.157; 6.U.157.166;
6.U.157.169; 6.U.157.172; 6.U.157.175; 6.U.157.240; 6.U.157.244;
6.U.166.228; 6.U.166.229; 6.U.166.230; 6.U.166.231; 6.U.166.236;
5 6.U.166.237; 6.U.166.238; 6.U.166.239; 6.U.166.154; 6.U.166.157;
6.U.166.166; 6.U.166.169; 6.U.166.172; 6.U.166.175; 6.U.166.240;
6.U.166.244; 6.U.169.228; 6.U.169.229; 6.U.169.230; 6.U.169.231;
6.U.169.236; 6.U.169.237; 6.U.169.238; 6.U.169.239; 6.U.169.154;
6.U.169.157; 6.U.169.166; 6.U.169.169; 6.U.169.172; 6.U.169.175;
10 6.U.169.240; 6.U.169.244; 6.U.172.228; 6.U.172.229; 6.U.172.230;
6.U.172.231; 6.U.172.236; 6.U.172.237; 6.U.172.238; 6.U.172.239;
6.U.172.154; 6.U.172.157; 6.U.172.166; 6.U.172.169; 6.U.172.172;
6.U.172.175; 6.U.172.240; 6.U.172.244; 6.U.175.228; 6.U.175.229;
6.U.175.230; 6.U.175.231; 6.U.175.236; 6.U.175.237; 6.U.175.238;
15 6.U.175.239; 6.U.175.154; 6.U.175.157; 6.U.175.166; 6.U.175.169;
6.U.175.172; 6.U.175.175; 6.U.175.240; 6.U.175.244; 6.U.240.228;
6.U.240.229; 6.U.240.230; 6.U.240.231; 6.U.240.236; 6.U.240.237;
6.U.240.238; 6.U.240.239; 6.U.240.154; 6.U.240.157; 6.U.240.166;
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20 6.U.244.228; 6.U.244.229; 6.U.244.230; 6.U.244.231; 6.U.244.236;
6.U.244.237; 6.U.244.238; 6.U.244.239; 6.U.244.154; 6.U.244.157;
6.U.244.166; 6.U.244.169; 6.U.244.172; 6.U.244.175; 6.U.244.240;
6.U.244.244;
- 25 Prodrugs of 6.W
6.W.228.228; 6.W.228.229; 6.W.228.230; 6.W.228.231; 6.W.228.236;
6.W.228.237; 6.W.228.238; 6.W.228.239; 6.W.228.154; 6.W.228.157;
6.W.228.166; 6.W.228.169; 6.W.228.172; 6.W.228.175; 6.W.228.240;
6.W.228.244; 6.W.229.228; 6.W.229.229; 6.W.229.230; 6.W.229.231;
30 6.W.229.236; 6.W.229.237; 6.W.229.238; 6.W.229.239; 6.W.229.154;

6.W.229.157; 6.W.229.166; 6.W.229.169; 6.W.229.172; 6.W.229.175;
6.W.229.240; 6.W.229.244; 6.W.230.228; 6.W.230.229; 6.W.230.230;
6.W.230.231; 6.W.230.236; 6.W.230.237; 6.W.230.238; 6.W.230.239;
6.W.230.154; 6.W.230.157; 6.W.230.166; 6.W.230.169; 6.W.230.172;
5 6.W.230.175; 6.W.230.240; 6.W.230.244; 6.W.231.228; 6.W.231.229;
6.W.231.230; 6.W.231.231; 6.W.231.236; 6.W.231.237; 6.W.231.238;
6.W.231.239; 6.W.231.154; 6.W.231.157; 6.W.231.166; 6.W.231.169;
6.W.231.172; 6.W.231.175; 6.W.231.240; 6.W.231.244; 6.W.236.228;
6.W.236.229; 6.W.236.230; 6.W.236.231; 6.W.236.236; 6.W.236.237;
10 6.W.236.238; 6.W.236.239; 6.W.236.154; 6.W.236.157; 6.W.236.166;
6.W.236.169; 6.W.236.172; 6.W.236.175; 6.W.236.240; 6.W.236.244;
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6.W.237.166; 6.W.237.169; 6.W.237.172; 6.W.237.175; 6.W.237.240;
15 6.W.237.244; 6.W.238.228; 6.W.238.229; 6.W.238.230; 6.W.238.231;
6.W.238.236; 6.W.238.237; 6.W.238.238; 6.W.238.239; 6.W.238.154;
6.W.238.157; 6.W.238.166; 6.W.238.169; 6.W.238.172; 6.W.238.175;
6.W.238.240; 6.W.238.244; 6.W.239.228; 6.W.239.229; 6.W.239.230;
6.W.239.231; 6.W.239.236; 6.W.239.237; 6.W.239.238; 6.W.239.239;
20 6.W.239.154; 6.W.239.157; 6.W.239.166; 6.W.239.169; 6.W.239.172;
6.W.239.175; 6.W.239.240; 6.W.239.244; 6.W.154.228; 6.W.154.229;
6.W.154.230; 6.W.154.231; 6.W.154.236; 6.W.154.237; 6.W.154.238;
6.W.154.239; 6.W.154.154; 6.W.154.157; 6.W.154.166; 6.W.154.169;
6.W.154.172; 6.W.154.175; 6.W.154.240; 6.W.154.244; 6.W.157.228;
25 6.W.157.229; 6.W.157.230; 6.W.157.231; 6.W.157.236; 6.W.157.237;
6.W.157.238; 6.W.157.239; 6.W.157.154; 6.W.157.157; 6.W.157.166;
6.W.157.169; 6.W.157.172; 6.W.157.175; 6.W.157.240; 6.W.157.244;
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6.W.166.237; 6.W.166.238; 6.W.166.239; 6.W.166.154; 6.W.166.157;
30 6.W.166.166; 6.W.166.169; 6.W.166.172; 6.W.166.175; 6.W.166.240;

6.W.166.244; 6.W.169.228; 6.W.169.229; 6.W.169.230; 6.W.169.231;
6.W.169.236; 6.W.169.237; 6.W.169.238; 6.W.169.239; 6.W.169.154;
6.W.169.157; 6.W.169.166; 6.W.169.169; 6.W.169.172; 6.W.169.175;
6.W.169.240; 6.W.169.244; 6.W.172.228; 6.W.172.229; 6.W.172.230;
5 6.W.172.231; 6.W.172.236; 6.W.172.237; 6.W.172.238; 6.W.172.239;
6.W.172.154; 6.W.172.157; 6.W.172.166; 6.W.172.169; 6.W.172.172;
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6.W.175.239; 6.W.175.154; 6.W.175.157; 6.W.175.166; 6.W.175.169;
10 6.W.175.172; 6.W.175.175; 6.W.175.240; 6.W.175.244; 6.W.240.228;
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15 6.W.244.237; 6.W.244.238; 6.W.244.239; 6.W.244.154; 6.W.244.157;
6.W.244.166; 6.W.244.169; 6.W.244.172; 6.W.244.175; 6.W.244.240;
6.W.244.244;

Prodrugs of 6.Y

20 6.Y.228.228; 6.Y.228.229; 6.Y.228.230; 6.Y.228.231; 6.Y.228.236;
6.Y.228.237; 6.Y.228.238; 6.Y.228.239; 6.Y.228.154; 6.Y.228.157;
6.Y.228.166; 6.Y.228.169; 6.Y.228.172; 6.Y.228.175; 6.Y.228.240;
6.Y.228.244; 6.Y.229.228; 6.Y.229.229; 6.Y.229.230; 6.Y.229.231;
6.Y.229.236; 6.Y.229.237; 6.Y.229.238; 6.Y.229.239; 6.Y.229.154;
25 6.Y.229.157; 6.Y.229.166; 6.Y.229.169; 6.Y.229.172; 6.Y.229.175;
6.Y.229.240; 6.Y.229.244; 6.Y.230.228; 6.Y.230.229; 6.Y.230.230;
6.Y.230.231; 6.Y.230.236; 6.Y.230.237; 6.Y.230.238; 6.Y.230.239;
6.Y.230.154; 6.Y.230.157; 6.Y.230.166; 6.Y.230.169; 6.Y.230.172;
6.Y.230.175; 6.Y.230.240; 6.Y.230.244; 6.Y.231.228; 6.Y.231.229;
30 6.Y.231.230; 6.Y.231.231; 6.Y.231.236; 6.Y.231.237; 6.Y.231.238;
6.Y.231.239; 6.Y.231.154; 6.Y.231.157; 6.Y.231.166; 6.Y.231.169;

6.Y.231.172; 6.Y.231.175; 6.Y.231.240; 6.Y.231.244; 6.Y.236.228;
6.Y.236.229; 6.Y.236.230; 6.Y.236.231; 6.Y.236.236; 6.Y.236.237;
6.Y.236.238; 6.Y.236.239; 6.Y.236.154; 6.Y.236.157; 6.Y.236.166;
6.Y.236.169; 6.Y.236.172; 6.Y.236.175; 6.Y.236.240; 6.Y.236.244;
5 6.Y.237.228; 6.Y.237.229; 6.Y.237.230; 6.Y.237.231; 6.Y.237.236;
6.Y.237.237; 6.Y.237.238; 6.Y.237.239; 6.Y.237.154; 6.Y.237.157;
6.Y.237.166; 6.Y.237.169; 6.Y.237.172; 6.Y.237.175; 6.Y.237.240;
6.Y.237.244; 6.Y.238.228; 6.Y.238.229; 6.Y.238.230; 6.Y.238.231;
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10 6.Y.238.157; 6.Y.238.166; 6.Y.238.169; 6.Y.238.172; 6.Y.238.175;
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Prodrugs of 7.AH

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Prodrugs of 7.AI

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Prodrugs of 7.AN

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Prodrugs of 7.AP

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- 30 Prodrugs of 7.AZ
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Prodrugs of 7.BF

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7.BF.247.223; 7.BF.247.240; 7.BF.247.244; 7.BF.247.243; 7.BF.247.247;

Prodrugs of 7.CI

25 7.CI.4.157; 7.CI.4.158; 7.CI.4.196; 7.CI.4.223; 7.CI.4.240; 7.CI.4.244;
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7.Cl.247.244; 7.Cl.247.243; 7.Cl.247.247;

Prodrugs of 7.CO

20 7.CO.4.157; 7.CO.4.158; 7.CO.4.196; 7.CO.4.223; 7.CO.4.240; 7.CO.4.244;
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25 7.CO.15.244; 7.CO.15.243; 7.CO.15.247; 7.CO.16.157; 7.CO.16.158;
7.CO.16.196; 7.CO.16.223; 7.CO.16.240; 7.CO.16.244; 7.CO.16.243;
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7.CO.26.158; 7.CO.26.196; 7.CO.26.223; 7.CO.26.240; 7.CO.26.244;
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10 7.CO.223.247; 7.CO.240.157; 7.CO.240.158; 7.CO.240.196; 7.CO.240.223;
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7.CO.4.223; 7.CO.4.240; 7.CO.4.244; 7.CO.4.243; 7.CO.4.247;

15

Prodrugs of 8.AH

- 8.AH.4.157; 8.AH.4.158; 8.AH.4.196; 8.AH.4.223; 8.AH.4.240; 8.AH.4.244;
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8.AH.5.240; 8.AH.5.244; 8.AH.5.243; 8.AH.5.247; 8.AH.7.157; 8.AH.7.158;
20 8.AH.7.196; 8.AH.7.223; 8.AH.7.240; 8.AH.7.244; 8.AH.7.243; 8.AH.7.247;
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25 8.AH.18.240; 8.AH.18.244; 8.AH.18.243; 8.AH.18.247; 8.AH.26.157;
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Prodrugs of 8.AJ

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Prodrugs of 8.AN

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 25 8.AN.55.240; 8.AN.55.244; 8.AN.55.243; 8.AN.55.247; 8.AN.56.157;
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5 Prodrugs of 8.AP

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10 8.AP.15.157; 8.AP.15.158; 8.AP.15.196; 8.AP.15.223; 8.AP.15.240;
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8.AP.18.240; 8.AP.18.244; 8.AP.18.243; 8.AP.18.247; 8.AP.26.157;
15 8.AP.26.158; 8.AP.26.196; 8.AP.26.223; 8.AP.26.240; 8.AP.26.244;
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Prodrugs of 8.AZ

- 8.AZ.4.157; 8.AZ.4.158; 8.AZ.4.196; 8.AZ.4.223; 8.AZ.4.240; 8.AZ.4.244;
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Prodrugs of 8.BF

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30 Prodrugs of 8.CI

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- 8.Cl.7.196; 8.Cl.7.223; 8.Cl.7.240; 8.Cl.7.244; 8.Cl.7.243; 8.Cl.7.247;
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25 Prodrugs of 8.CO

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Prodrugs of 9.AH

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Prodrugs of 9.AI

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Prodrugs of 9.AN

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Prodrugs of 9.AP

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Prodrugs of 9.AZ

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5 Prodrugs of 9.BF

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Prodrugs of 9.Cl

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30 Prodrugs of 9.CO

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Prodrugs of 10.AH

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Prodrugs of 10.AI

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Prodrugs of 11.AJ

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Prodrugs of 11.AN

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25 Prodrugs of 11.AP

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25 Prodrugs of 11.AZ

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25 Prodrugs of 11.BF

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12.BF.15.243; 12.BF.15.247; 12.BF.16.157; 12.BF.16.158; 12.BF.16.196;
30 12.BF.16.223; 12.BF.16.240; 12.BF.16.244; 12.BF.16.243; 12.BF.16.247;
12.BF.18.157; 12.BF.18.158; 12.BF.18.196; 12.BF.18.223; 12.BF.18.240;
12.BF.18.244; 12.BF.18.243; 12.BF.18.247; 12.BF.26.157; 12.BF.26.158;
12.BF.26.196; 12.BF.26.223; 12.BF.26.240; 12.BF.26.244; 12.BF.26.243;

12.BF.26.247; 12.BF.27.157; 12.BF.27.158; 12.BF.27.196; 12.BF.27.223;
12.BF.27.240; 12.BF.27.244; 12.BF.27.243; 12.BF.27.247; 12.BF.29.157;
12.BF.29.158; 12.BF.29.196; 12.BF.29.223; 12.BF.29.240; 12.BF.29.244;
12.BF.29.243; 12.BF.29.247; 12.BF.54.157; 12.BF.54.158; 12.BF.54.196;
5 12.BF.54.223; 12.BF.54.240; 12.BF.54.244; 12.BF.54.243; 12.BF.54.247;
12.BF.55.157; 12.BF.55.158; 12.BF.55.196; 12.BF.55.223; 12.BF.55.240;
12.BF.55.244; 12.BF.55.243; 12.BF.55.247; 12.BF.56.157; 12.BF.56.158;
12.BF.56.196; 12.BF.56.223; 12.BF.56.240; 12.BF.56.244; 12.BF.56.243;
12.BF.56.247; 12.BF.157.157; 12.BF.157.158; 12.BF.157.196; 12.BF.157.223;
10 12.BF.157.240; 12.BF.157.244; 12.BF.157.243; 12.BF.157.247; 12.BF.196.157;
12.BF.196.158; 12.BF.196.196; 12.BF.196.223; 12.BF.196.240; 12.BF.196.244;
12.BF.196.243; 12.BF.196.247; 12.BF.223.157; 12.BF.223.158; 12.BF.223.196;
12.BF.223.223; 12.BF.223.240; 12.BF.223.244; 12.BF.223.243; 12.BF.223.247;
12.BF.240.157; 12.BF.240.158; 12.BF.240.196; 12.BF.240.223; 12.BF.240.240;
15 12.BF.240.244; 12.BF.240.243; 12.BF.240.247; 12.BF.244.157; 12.BF.244.158;
12.BF.244.196; 12.BF.244.223; 12.BF.244.240; 12.BF.244.244; 12.BF.244.243;
12.BF.244.247; 12.BF.247.157; 12.BF.247.158; 12.BF.247.196; 12.BF.247.223;
12.BF.247.240; 12.BF.247.244; 12.BF.247.243; 12.BF.247.247;

20 Prodrugs of 12.CI

12.CI.4.157; 12.CI.4.158; 12.CI.4.196; 12.CI.4.223; 12.CI.4.240;
12.CI.4.244; 12.CI.4.243; 12.CI.4.247; 12.CI.5.157; 12.CI.5.158; 12.CI.5.196;
12.CI.5.223; 12.CI.5.240; 12.CI.5.244; 12.CI.5.243; 12.CI.5.247; 12.CI.7.157;
12.CI.7.158; 12.CI.7.196; 12.CI.7.223; 12.CI.7.240; 12.CI.7.244; 12.CI.7.243;
25 12.CI.7.247; 12.CI.15.157; 12.CI.15.158; 12.CI.15.196; 12.CI.15.223;
12.CI.15.240; 12.CI.15.244; 12.CI.15.243; 12.CI.15.247; 12.CI.16.157;
12.CI.16.158; 12.CI.16.196; 12.CI.16.223; 12.CI.16.240; 12.CI.16.244;
12.CI.16.243; 12.CI.16.247; 12.CI.18.157; 12.CI.18.158; 12.CI.18.196;
12.CI.18.223; 12.CI.18.240; 12.CI.18.244; 12.CI.18.243; 12.CI.18.247;
30 12.CI.26.157; 12.CI.26.158; 12.CI.26.196; 12.CI.26.223; 12.CI.26.240;
12.CI.26.244; 12.CI.26.243; 12.CI.26.247; 12.CI.27.157; 12.CI.27.158;
12.CI.27.196; 12.CI.27.223; 12.CI.27.240; 12.CI.27.244; 12.CI.27.243;
12.CI.27.247; 12.CI.29.157; 12.CI.29.158; 12.CI.29.196; 12.CI.29.223;

- 12.CI.29.240; 12.CI.29.244; 12.CI.29.243; 12.CI.29.247; 12.CI.54.157;
12.CI.54.158; 12.CI.54.196; 12.CI.54.223; 12.CI.54.240; 12.CI.54.244;
12.CI.54.243; 12.CI.54.247; 12.CI.55.157; 12.CI.55.158; 12.CI.55.196;
12.CI.55.223; 12.CI.55.240; 12.CI.55.244; 12.CI.55.243; 12.CI.55.247;
5 12.CI.56.157; 12.CI.56.158; 12.CI.56.196; 12.CI.56.223; 12.CI.56.240;
12.CI.56.244; 12.CI.56.243; 12.CI.56.247; 12.CI.157.157; 12.CI.157.158;
12.CI.157.196; 12.CI.157.223; 12.CI.157.240; 12.CI.157.244; 12.CI.157.243;
12.CI.157.247; 12.CI.196.157; 12.CI.196.158; 12.CI.196.196; 12.CI.196.223;
12.CI.196.240; 12.CI.196.244; 12.CI.196.243; 12.CI.196.247; 12.CI.223.157;
10 12.CI.223.158; 12.CI.223.196; 12.CI.223.223; 12.CI.223.240; 12.CI.223.244;
12.CI.223.243; 12.CI.223.247; 12.CI.240.157; 12.CI.240.158; 12.CI.240.196;
12.CI.240.223; 12.CI.240.240; 12.CI.240.244; 12.CI.240.243; 12.CI.240.247;
12.CI.244.157; 12.CI.244.158; 12.CI.244.196; 12.CI.244.223; 12.CI.244.240;
12.CI.244.244; 12.CI.244.243; 12.CI.244.247; 12.CI.247.157; 12.CI.247.158;
15 12.CI.247.196; 12.CI.247.223; 12.CI.247.240; 12.CI.247.244; 12.CI.247.243;
12.CI.247.247;

Prodrugs of 12.CO

- 12.CO.4.157; 12.CO.4.158; 12.CO.4.196; 12.CO.4.223; 12.CO.4.240;
20 12.CO.4.244; 12.CO.4.243; 12.CO.4.247; 12.CO.5.157; 12.CO.5.158;
12.CO.5.196; 12.CO.5.223; 12.CO.5.240; 12.CO.5.244; 12.CO.5.243;
12.CO.5.247; 12.CO.7.157; 12.CO.7.158; 12.CO.7.196; 12.CO.7.223;
12.CO.7.240; 12.CO.7.244; 12.CO.7.243; 12.CO.7.247; 12.CO.15.157;
12.CO.15.158; 12.CO.15.196; 12.CO.15.223; 12.CO.15.240; 12.CO.15.244;
25 12.CO.15.243; 12.CO.15.247; 12.CO.16.157; 12.CO.16.158; 12.CO.16.196;
12.CO.16.223; 12.CO.16.240; 12.CO.16.244; 12.CO.16.243; 12.CO.16.247;
12.CO.18.157; 12.CO.18.158; 12.CO.18.196; 12.CO.18.223; 12.CO.18.240;
12.CO.18.244; 12.CO.18.243; 12.CO.18.247; 12.CO.26.157; 12.CO.26.158;
12.CO.26.196; 12.CO.26.223; 12.CO.26.240; 12.CO.26.244; 12.CO.26.243;
30 12.CO.26.247; 12.CO.27.157; 12.CO.27.158; 12.CO.27.196; 12.CO.27.223;
12.CO.27.240; 12.CO.27.244; 12.CO.27.243; 12.CO.27.247; 12.CO.29.157;
12.CO.29.158; 12.CO.29.196; 12.CO.29.223; 12.CO.29.240; 12.CO.29.244;
12.CO.29.243; 12.CO.29.247; 12.CO.54.157; 12.CO.54.158; 12.CO.54.196;

- 12.CO.54.223; 12.CO.54.240; 12.CO.54.244; 12.CO.54.243; 12.CO.54.247;
12.CO.55.157; 12.CO.55.158; 12.CO.55.196; 12.CO.55.223; 12.CO.55.240;
12.CO.55.244; 12.CO.55.243; 12.CO.55.247; 12.CO.56.157; 12.CO.56.158;
12.CO.56.196; 12.CO.56.223; 12.CO.56.240; 12.CO.56.244; 12.CO.56.243;
5 12.CO.56.247; 12.CO.157.157; 12.CO.157.158; 12.CO.157.196;
12.CO.157.223; 12.CO.157.240; 12.CO.157.244; 12.CO.157.243;
12.CO.157.247; 12.CO.196.157; 12.CO.196.158; 12.CO.196.196;
12.CO.196.223; 12.CO.196.240; 12.CO.196.244; 12.CO.196.243;
12.CO.196.247; 12.CO.223.157; 12.CO.223.158; 12.CO.223.196;
10 12.CO.223.223; 12.CO.223.240; 12.CO.223.244; 12.CO.223.243;
12.CO.223.247; 12.CO.240.157; 12.CO.240.158; 12.CO.240.196;
12.CO.240.223; 12.CO.240.240; 12.CO.240.244; 12.CO.240.243;
12.CO.240.247; 12.CO.244.157; 12.CO.244.158; 12.CO.244.196;
12.CO.244.223; 12.CO.244.240; 12.CO.244.244; 12.CO.244.243;
15 12.CO.244.247; 12.CO.247.157; 12.CO.247.158; 12.CO.247.196;
12.CO.247.223; 12.CO.247.240; 12.CO.247.244; 12.CO.247.243;
12.CO.247.247.

Prodrugs of 13.B

- 20 13.B.228.228; 13.B.228.229; 13.B.228.230; 13.B.228.231; 13.B.228.236;
13.B.228.237; 13.B.228.238; 13.B.228.239; 13.B.228.154; 13.B.228.157;
13.B.228.166; 13.B.228.169; 13.B.228.172; 13.B.228.175; 13.B.228.240;
13.B.228.244; 13.B.229.228; 13.B.229.229; 13.B.229.230; 13.B.229.231;
13.B.229.236; 13.B.229.237; 13.B.229.238; 13.B.229.239; 13.B.229.154;
25 13.B.229.157; 13.B.229.166; 13.B.229.169; 13.B.229.172; 13.B.229.175;
13.B.229.240; 13.B.229.244; 13.B.230.228; 13.B.230.229; 13.B.230.230;
13.B.230.231; 13.B.230.236; 13.B.230.237; 13.B.230.238; 13.B.230.239;
13.B.230.154; 13.B.230.157; 13.B.230.166; 13.B.230.169; 13.B.230.172;
13.B.230.175; 13.B.230.240; 13.B.230.244; 13.B.231.228; 13.B.231.229;
30 13.B.231.230; 13.B.231.231; 13.B.231.236; 13.B.231.237; 13.B.231.238;
13.B.231.239; 13.B.231.154; 13.B.231.157; 13.B.231.166; 13.B.231.169;

13.B.231.172; 13.B.231.175; 13.B.231.240; 13.B.231.244; 13.B.236.228;
13.B.236.229; 13.B.236.230; 13.B.236.231; 13.B.236.236; 13.B.236.237;
13.B.236.238; 13.B.236.239; 13.B.236.154; 13.B.236.157; 13.B.236.166;
13.B.236.169; 13.B.236.172; 13.B.236.175; 13.B.236.240; 13.B.236.244;
5 13.B.237.228; 13.B.237.229; 13.B.237.230; 13.B.237.231; 13.B.237.236;
13.B.237.237; 13.B.237.238; 13.B.237.239; 13.B.237.154; 13.B.237.157;
13.B.237.166; 13.B.237.169; 13.B.237.172; 13.B.237.175; 13.B.237.240;
13.B.237.244; 13.B.238.228; 13.B.238.229; 13.B.238.230; 13.B.238.231;
13.B.238.236; 13.B.238.237; 13.B.238.238; 13.B.238.239; 13.B.238.154;
10 13.B.238.157; 13.B.238.166; 13.B.238.169; 13.B.238.172; 13.B.238.175;
13.B.238.240; 13.B.238.244; 13.B.239.228; 13.B.239.229; 13.B.239.230;
13.B.239.231; 13.B.239.236; 13.B.239.237; 13.B.239.238; 13.B.239.239;
13.B.239.154; 13.B.239.157; 13.B.239.166; 13.B.239.169; 13.B.239.172;
13.B.239.175; 13.B.239.240; 13.B.239.244; 13.B.154.228; 13.B.154.229;
15 13.B.154.230; 13.B.154.231; 13.B.154.236; 13.B.154.237; 13.B.154.238;
13.B.154.239; 13.B.154.154; 13.B.154.157; 13.B.154.166; 13.B.154.169;
13.B.154.172; 13.B.154.175; 13.B.154.240; 13.B.154.244; 13.B.157.228;
13.B.157.229; 13.B.157.230; 13.B.157.231; 13.B.157.236; 13.B.157.237;
13.B.157.238; 13.B.157.239; 13.B.157.154; 13.B.157.157; 13.B.157.166;
20 13.B.157.169; 13.B.157.172; 13.B.157.175; 13.B.157.240; 13.B.157.244;
13.B.166.228; 13.B.166.229; 13.B.166.230; 13.B.166.231; 13.B.166.236;
13.B.166.237; 13.B.166.238; 13.B.166.239; 13.B.166.154; 13.B.166.157;
13.B.166.166; 13.B.166.169; 13.B.166.172; 13.B.166.175; 13.B.166.240;
13.B.166.244; 13.B.169.228; 13.B.169.229; 13.B.169.230; 13.B.169.231;
25 13.B.169.236; 13.B.169.237; 13.B.169.238; 13.B.169.239; 13.B.169.154;
13.B.169.157; 13.B.169.166; 13.B.169.169; 13.B.169.172; 13.B.169.175;
13.B.169.240; 13.B.169.244; 13.B.172.228; 13.B.172.229; 13.B.172.230;
13.B.172.231; 13.B.172.236; 13.B.172.237; 13.B.172.238; 13.B.172.239;
13.B.172.154; 13.B.172.157; 13.B.172.166; 13.B.172.169; 13.B.172.172;
30 13.B.172.175; 13.B.172.240; 13.B.172.244; 13.B.175.228; 13.B.175.229;

13.B.175.230; 13.B.175.231; 13.B.175.236; 13.B.175.237; 13.B.175.238;
13.B.175.239; 13.B.175.154; 13.B.175.157; 13.B.175.166; 13.B.175.169;
13.B.175.172; 13.B.175.175; 13.B.175.240; 13.B.175.244; 13.B.240.228;
13.B.240.229; 13.B.240.230; 13.B.240.231; 13.B.240.236; 13.B.240.237;
5 13.B.240.238; 13.B.240.239; 13.B.240.154; 13.B.240.157; 13.B.240.166;
13.B.240.169; 13.B.240.172; 13.B.240.175; 13.B.240.240; 13.B.240.244;
13.B.244.228; 13.B.244.229; 13.B.244.230; 13.B.244.231; 13.B.244.236;
13.B.244.237; 13.B.244.238; 13.B.244.239; 13.B.244.154; 13.B.244.157;
13.B.244.166; 13.B.244.169; 13.B.244.172; 13.B.244.175; 13.B.244.240;
10 13.B.244.244;

Prodrugs of 13.D

13.D.228.228; 13.D.228.229; 13.D.228.230; 13.D.228.231; 13.D.228.236;
13.D.228.237; 13.D.228.238; 13.D.228.239; 13.D.228.154; 13.D.228.157;
15 13.D.228.166; 13.D.228.169; 13.D.228.172; 13.D.228.175; 13.D.228.240;
13.D.228.244; 13.D.229.228; 13.D.229.229; 13.D.229.230; 13.D.229.231;
13.D.229.236; 13.D.229.237; 13.D.229.238; 13.D.229.239; 13.D.229.154;
13.D.229.157; 13.D.229.166; 13.D.229.169; 13.D.229.172; 13.D.229.175;
13.D.229.240; 13.D.229.244; 13.D.230.228; 13.D.230.229; 13.D.230.230;
20 13.D.230.231; 13.D.230.236; 13.D.230.237; 13.D.230.238; 13.D.230.239;
13.D.230.154; 13.D.230.157; 13.D.230.166; 13.D.230.169; 13.D.230.172;
13.D.230.175; 13.D.230.240; 13.D.230.244; 13.D.231.228; 13.D.231.229;
13.D.231.230; 13.D.231.231; 13.D.231.236; 13.D.231.237; 13.D.231.238;
13.D.231.239; 13.D.231.154; 13.D.231.157; 13.D.231.166; 13.D.231.169;
25 13.D.231.172; 13.D.231.175; 13.D.231.240; 13.D.231.244; 13.D.236.228;
13.D.236.229; 13.D.236.230; 13.D.236.231; 13.D.236.236; 13.D.236.237;
13.D.236.238; 13.D.236.239; 13.D.236.154; 13.D.236.157; 13.D.236.166;
13.D.236.169; 13.D.236.172; 13.D.236.175; 13.D.236.240; 13.D.236.244;
13.D.237.228; 13.D.237.229; 13.D.237.230; 13.D.237.231; 13.D.237.236;
30 13.D.237.237; 13.D.237.238; 13.D.237.239; 13.D.237.154; 13.D.237.157;

13.D.237.166; 13.D.237.169; 13.D.237.172; 13.D.237.175; 13.D.237.240;
13.D.237.244; 13.D.238.228; 13.D.238.229; 13.D.238.230; 13.D.238.231;
13.D.238.236; 13.D.238.237; 13.D.238.238; 13.D.238.239; 13.D.238.154;
13.D.238.157; 13.D.238.166; 13.D.238.169; 13.D.238.172; 13.D.238.175;
5 13.D.238.240; 13.D.238.244; 13.D.239.228; 13.D.239.229; 13.D.239.230;
13.D.239.231; 13.D.239.236; 13.D.239.237; 13.D.239.238; 13.D.239.239;
13.D.239.154; 13.D.239.157; 13.D.239.166; 13.D.239.169; 13.D.239.172;
13.D.239.175; 13.D.239.240; 13.D.239.244; 13.D.154.228; 13.D.154.229;
13.D.154.230; 13.D.154.231; 13.D.154.236; 13.D.154.237; 13.D.154.238;
10 13.D.154.239; 13.D.154.154; 13.D.154.157; 13.D.154.166; 13.D.154.169;
13.D.154.172; 13.D.154.175; 13.D.154.240; 13.D.154.244; 13.D.157.228;
13.D.157.229; 13.D.157.230; 13.D.157.231; 13.D.157.236; 13.D.157.237;
13.D.157.238; 13.D.157.239; 13.D.157.154; 13.D.157.157; 13.D.157.166;
13.D.157.169; 13.D.157.172; 13.D.157.175; 13.D.157.240; 13.D.157.244;
15 13.D.166.228; 13.D.166.229; 13.D.166.230; 13.D.166.231; 13.D.166.236;
13.D.166.237; 13.D.166.238; 13.D.166.239; 13.D.166.154; 13.D.166.157;
13.D.166.166; 13.D.166.169; 13.D.166.172; 13.D.166.175; 13.D.166.240;
13.D.166.244; 13.D.169.228; 13.D.169.229; 13.D.169.230; 13.D.169.231;
13.D.169.236; 13.D.169.237; 13.D.169.238; 13.D.169.239; 13.D.169.154;
20 13.D.169.157; 13.D.169.166; 13.D.169.169; 13.D.169.172; 13.D.169.175;
13.D.169.240; 13.D.169.244; 13.D.172.228; 13.D.172.229; 13.D.172.230;
13.D.172.231; 13.D.172.236; 13.D.172.237; 13.D.172.238; 13.D.172.239;
13.D.172.154; 13.D.172.157; 13.D.172.166; 13.D.172.169; 13.D.172.172;
13.D.172.175; 13.D.172.240; 13.D.172.244; 13.D.175.228; 13.D.175.229;
25 13.D.175.230; 13.D.175.231; 13.D.175.236; 13.D.175.237; 13.D.175.238;
13.D.175.239; 13.D.175.154; 13.D.175.157; 13.D.175.166; 13.D.175.169;
13.D.175.172; 13.D.175.175; 13.D.175.240; 13.D.175.244; 13.D.240.228;
13.D.240.229; 13.D.240.230; 13.D.240.231; 13.D.240.236; 13.D.240.237;
13.D.240.238; 13.D.240.239; 13.D.240.154; 13.D.240.157; 13.D.240.166;
30 13.D.240.169; 13.D.240.172; 13.D.240.175; 13.D.240.240; 13.D.240.244;

13.D.244.228; 13.D.244.229; 13.D.244.230; 13.D.244.231; 13.D.244.236;
13.D.244.237; 13.D.244.238; 13.D.244.239; 13.D.244.154; 13.D.244.157;
13.D.244.166; 13.D.244.169; 13.D.244.172; 13.D.244.175; 13.D.244.240;
13.D.244.244;

5

Prodrugs of 13.E

13.E.228.228; 13.E.228.229; 13.E.228.230; 13.E.228.231; 13.E.228.236;
13.E.228.237; 13.E.228.238; 13.E.228.239; 13.E.228.154; 13.E.228.157;
13.E.228.166; 13.E.228.169; 13.E.228.172; 13.E.228.175; 13.E.228.240;
10 13.E.228.244; 13.E.229.228; 13.E.229.229; 13.E.229.230; 13.E.229.231;
13.E.229.236; 13.E.229.237; 13.E.229.238; 13.E.229.239; 13.E.229.154;
13.E.229.157; 13.E.229.166; 13.E.229.169; 13.E.229.172; 13.E.229.175;
13.E.229.240; 13.E.229.244; 13.E.230.228; 13.E.230.229; 13.E.230.230;
13.E.230.231; 13.E.230.236; 13.E.230.237; 13.E.230.238; 13.E.230.239;
15 13.E.230.154; 13.E.230.157; 13.E.230.166; 13.E.230.169; 13.E.230.172;
13.E.230.175; 13.E.230.240; 13.E.230.244; 13.E.231.228; 13.E.231.229;
13.E.231.230; 13.E.231.231; 13.E.231.236; 13.E.231.237; 13.E.231.238;
13.E.231.239; 13.E.231.154; 13.E.231.157; 13.E.231.166; 13.E.231.169;
13.E.231.172; 13.E.231.175; 13.E.231.240; 13.E.231.244; 13.E.236.228;
20 13.E.236.229; 13.E.236.230; 13.E.236.231; 13.E.236.236; 13.E.236.237;
13.E.236.238; 13.E.236.239; 13.E.236.154; 13.E.236.157; 13.E.236.166;
13.E.236.169; 13.E.236.172; 13.E.236.175; 13.E.236.240; 13.E.236.244;
13.E.237.228; 13.E.237.229; 13.E.237.230; 13.E.237.231; 13.E.237.236;
13.E.237.237; 13.E.237.238; 13.E.237.239; 13.E.237.154; 13.E.237.157;
25 13.E.237.166; 13.E.237.169; 13.E.237.172; 13.E.237.175; 13.E.237.240;
13.E.237.244; 13.E.238.228; 13.E.238.229; 13.E.238.230; 13.E.238.231;
13.E.238.236; 13.E.238.237; 13.E.238.238; 13.E.238.239; 13.E.238.154;
13.E.238.157; 13.E.238.166; 13.E.238.169; 13.E.238.172; 13.E.238.175;
13.E.238.240; 13.E.238.244; 13.E.239.228; 13.E.239.229; 13.E.239.230;
30 13.E.239.231; 13.E.239.236; 13.E.239.237; 13.E.239.238; 13.E.239.239;

13.E.239.154; 13.E.239.157; 13.E.239.166; 13.E.239.169; 13.E.239.172;
13.E.239.175; 13.E.239.240; 13.E.239.244; 13.E.154.228; 13.E.154.229;
13.E.154.230; 13.E.154.231; 13.E.154.236; 13.E.154.237; 13.E.154.238;
13.E.154.239; 13.E.154.154; 13.E.154.157; 13.E.154.166; 13.E.154.169;
5 13.E.154.172; 13.E.154.175; 13.E.154.240; 13.E.154.244; 13.E.157.228;
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13.E.157.238; 13.E.157.239; 13.E.157.154; 13.E.157.157; 13.E.157.166;
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13.E.166.228; 13.E.166.229; 13.E.166.230; 13.E.166.231; 13.E.166.236;
10 13.E.166.237; 13.E.166.238; 13.E.166.239; 13.E.166.154; 13.E.166.157;
13.E.166.166; 13.E.166.169; 13.E.166.172; 13.E.166.175; 13.E.166.240;
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13.E.169.157; 13.E.169.166; 13.E.169.169; 13.E.169.172; 13.E.169.175;
15 13.E.169.240; 13.E.169.244; 13.E.172.228; 13.E.172.229; 13.E.172.230;
13.E.172.231; 13.E.172.236; 13.E.172.237; 13.E.172.238; 13.E.172.239;
13.E.172.154; 13.E.172.157; 13.E.172.166; 13.E.172.169; 13.E.172.172;
13.E.172.175; 13.E.172.240; 13.E.172.244; 13.E.175.228; 13.E.175.229;
13.E.175.230; 13.E.175.231; 13.E.175.236; 13.E.175.237; 13.E.175.238;
20 13.E.175.239; 13.E.175.154; 13.E.175.157; 13.E.175.166; 13.E.175.169;
13.E.175.172; 13.E.175.175; 13.E.175.240; 13.E.175.244; 13.E.240.228;
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25 13.E.244.228; 13.E.244.229; 13.E.244.230; 13.E.244.231; 13.E.244.236;
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13.E.244.166; 13.E.244.169; 13.E.244.172; 13.E.244.175; 13.E.244.240;
13.E.244.244;

30 Prodrugs of 13.G

13.G.228.228; 13.G.228.229; 13.G.228.230; 13.G.228.231; 13.G.228.236;
13.G.228.237; 13.G.228.238; 13.G.228.239; 13.G.228.154; 13.G.228.157;
13.G.228.166; 13.G.228.169; 13.G.228.172; 13.G.228.175; 13.G.228.240;
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5 13.G.229.236; 13.G.229.237; 13.G.229.238; 13.G.229.239; 13.G.229.154;
13.G.229.157; 13.G.229.166; 13.G.229.169; 13.G.229.172; 13.G.229.175;
13.G.229.240; 13.G.229.244; 13.G.230.228; 13.G.230.229; 13.G.230.230;
13.G.230.231; 13.G.230.236; 13.G.230.237; 13.G.230.238; 13.G.230.239;
13.G.230.154; 13.G.230.157; 13.G.230.166; 13.G.230.169; 13.G.230.172;
10 13.G.230.175; 13.G.230.240; 13.G.230.244; 13.G.231.228; 13.G.231.229;
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13.G.231.239; 13.G.231.154; 13.G.231.157; 13.G.231.166; 13.G.231.169;
13.G.231.172; 13.G.231.175; 13.G.231.240; 13.G.231.244; 13.G.236.228;
13.G.236.229; 13.G.236.230; 13.G.236.231; 13.G.236.236; 13.G.236.237;
15 13.G.236.238; 13.G.236.239; 13.G.236.154; 13.G.236.157; 13.G.236.166;
13.G.236.169; 13.G.236.172; 13.G.236.175; 13.G.236.240; 13.G.236.244;
13.G.237.228; 13.G.237.229; 13.G.237.230; 13.G.237.231; 13.G.237.236;
13.G.237.237; 13.G.237.238; 13.G.237.239; 13.G.237.154; 13.G.237.157;
13.G.237.166; 13.G.237.169; 13.G.237.172; 13.G.237.175; 13.G.237.240;
20 13.G.237.244; 13.G.238.228; 13.G.238.229; 13.G.238.230; 13.G.238.231;
13.G.238.236; 13.G.238.237; 13.G.238.238; 13.G.238.239; 13.G.238.154;
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25 13.G.239.154; 13.G.239.157; 13.G.239.166; 13.G.239.169; 13.G.239.172;
13.G.239.175; 13.G.239.240; 13.G.239.244; 13.G.154.228; 13.G.154.229;
13.G.154.230; 13.G.154.231; 13.G.154.236; 13.G.154.237; 13.G.154.238;
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13.G.154.172; 13.G.154.175; 13.G.154.240; 13.G.154.244; 13.G.157.228;
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13.G.157.238; 13.G.157.239; 13.G.157.154; 13.G.157.157; 13.G.157.166;
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13.G.169.240; 13.G.169.244; 13.G.172.228; 13.G.172.229; 13.G.172.230;
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15 13.G.175.172; 13.G.175.175; 13.G.175.240; 13.G.175.244; 13.G.240.228;
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20 13.G.244.237; 13.G.244.238; 13.G.244.239; 13.G.244.154; 13.G.244.157;
13.G.244.166; 13.G.244.169; 13.G.244.172; 13.G.244.175; 13.G.244.240;
13.G.244.244;

Prodrugs of 13.I

25 13.I.228.228; 13.I.228.229; 13.I.228.230; 13.I.228.231; 13.I.228.236;
13.I.228.237; 13.I.228.238; 13.I.228.239; 13.I.228.154; 13.I.228.157;
13.I.228.166; 13.I.228.169; 13.I.228.172; 13.I.228.175; 13.I.228.240;
13.I.228.244; 13.I.229.228; 13.I.229.229; 13.I.229.230; 13.I.229.231;
13.I.229.236; 13.I.229.237; 13.I.229.238; 13.I.229.239; 13.I.229.154;
30 13.I.229.157; 13.I.229.166; 13.I.229.169; 13.I.229.172; 13.I.229.175;

13.I.229.240; 13.I.229.244; 13.I.230.228; 13.I.230.229; 13.I.230.230;
13.I.230.231; 13.I.230.236; 13.I.230.237; 13.I.230.238; 13.I.230.239;
13.I.230.154; 13.I.230.157; 13.I.230.166; 13.I.230.169; 13.I.230.172;
13.I.230.175; 13.I.230.240; 13.I.230.244; 13.I.231.228; 13.I.231.229;
5 13.I.231.230; 13.I.231.231; 13.I.231.236; 13.I.231.237; 13.I.231.238;
13.I.231.239; 13.I.231.154; 13.I.231.157; 13.I.231.166; 13.I.231.169;
13.I.231.172; 13.I.231.175; 13.I.231.240; 13.I.231.244; 13.I.236.228;
13.I.236.229; 13.I.236.230; 13.I.236.231; 13.I.236.236; 13.I.236.237;
13.I.236.238; 13.I.236.239; 13.I.236.154; 13.I.236.157; 13.I.236.166;
10 13.I.236.169; 13.I.236.172; 13.I.236.175; 13.I.236.240; 13.I.236.244;
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13.I.237.237; 13.I.237.238; 13.I.237.239; 13.I.237.154; 13.I.237.157;
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13.I.237.244; 13.I.238.228; 13.I.238.229; 13.I.238.230; 13.I.238.231;
15 13.I.238.236; 13.I.238.237; 13.I.238.238; 13.I.238.239; 13.I.238.154;
13.I.238.157; 13.I.238.166; 13.I.238.169; 13.I.238.172; 13.I.238.175;
13.I.238.240; 13.I.238.244; 13.I.239.228; 13.I.239.229; 13.I.239.230;
13.I.239.231; 13.I.239.236; 13.I.239.237; 13.I.239.238; 13.I.239.239;
13.I.239.154; 13.I.239.157; 13.I.239.166; 13.I.239.169; 13.I.239.172;
20 13.I.239.175; 13.I.239.240; 13.I.239.244; 13.I.154.228; 13.I.154.229;
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13.I.157.229; 13.I.157.230; 13.I.157.231; 13.I.157.236; 13.I.157.237;
25 13.I.157.238; 13.I.157.239; 13.I.157.154; 13.I.157.157; 13.I.157.166;
13.I.157.169; 13.I.157.172; 13.I.157.175; 13.I.157.240; 13.I.157.244;
13.I.166.228; 13.I.166.229; 13.I.166.230; 13.I.166.231; 13.I.166.236;
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13.I.166.166; 13.I.166.169; 13.I.166.172; 13.I.166.175; 13.I.166.240;
30 13.I.166.244; 13.I.169.228; 13.I.169.229; 13.I.169.230; 13.I.169.231;

13.I.169.236; 13.I.169.237; 13.I.169.238; 13.I.169.239; 13.I.169.154;
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13.I.172.231; 13.I.172.236; 13.I.172.237; 13.I.172.238; 13.I.172.239;
5 13.I.172.154; 13.I.172.157; 13.I.172.166; 13.I.172.169; 13.I.172.172;
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13.I.175.172; 13.I.175.175; 13.I.175.240; 13.I.175.244; 13.I.240.228;
10 13.I.240.229; 13.I.240.230; 13.I.240.231; 13.I.240.236; 13.I.240.237;
13.I.240.238; 13.I.240.239; 13.I.240.154; 13.I.240.157; 13.I.240.166;
13.I.240.169; 13.I.240.172; 13.I.240.175; 13.I.240.240; 13.I.240.244;
13.I.244.228; 13.I.244.229; 13.I.244.230; 13.I.244.231; 13.I.244.236;
13.I.244.237; 13.I.244.238; 13.I.244.239; 13.I.244.154; 13.I.244.157;
15 13.I.244.166; 13.I.244.169; 13.I.244.172; 13.I.244.175; 13.I.244.240;
13.I.244.244; .

Prodrugs of 13.I

13.J.228.228; 13.J.228.229; 13.J.228.230; 13.J.228.231; 13.J.228.236;
20 13.J.228.237; 13.J.228.238; 13.J.228.239; 13.J.228.154; 13.J.228.157;
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13.J.231.230; 13.J.231.231; 13.J.231.236; 13.J.231.237; 13.J.231.238;
30 13.J.231.239; 13.J.231.154; 13.J.231.157; 13.J.231.166; 13.J.231.169;

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5 13.J.237.228; 13.J.237.229; 13.J.237.230; 13.J.237.231; 13.J.237.236;
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10 13.J.238.157; 13.J.238.166; 13.J.238.169; 13.J.238.172; 13.J.238.175;
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25 13.J.169.236; 13.J.169.237; 13.J.169.238; 13.J.169.239; 13.J.169.154;
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Prodrugs of 13.W

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Prodrugs of 14.AH

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Prodrugs of 14.AJ

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Prodrugs of 14.AN

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5 All literature and patent citations above are hereby expressly
incorporated by reference at the locations of their citation. Specifically cited
sections or pages of the above cited works are incorporated by reference with
specificity. The invention has been described in detail sufficient to allow one of
ordinary skill in the art to make and use the subject matter of the following
Claims. It is apparent that certain modifications of the methods and
10 compositions of the following Claims can be made within the scope and spirit of
the invention.

In the claims hereinbelow, the subscript and superscripts of a given
variable are distinct. For example, R_1 is distinct from R^1 .

15

Claims

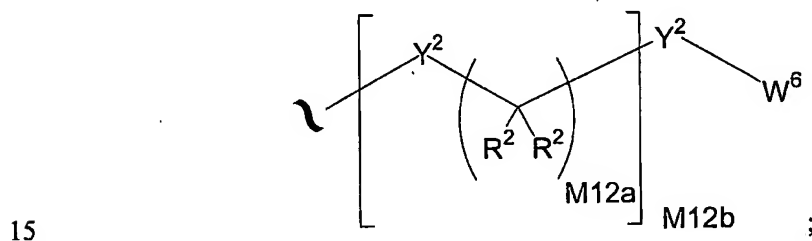
We claim:

1. A conjugate comprising an immuno-modulatory compound linked to one
5 or more phosphonate groups; or a pharmaceutically acceptable salt or solvate thereof.

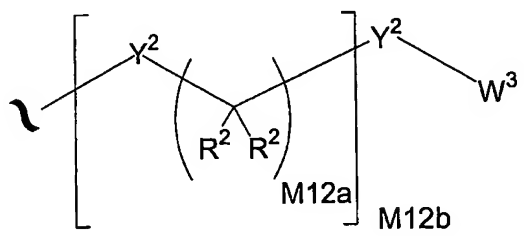
2. The conjugate of claim 1, or a pharmaceutically acceptable salt or solvate thereof, that is a compound of any one of formulae 500-547 substituted with one
10 or more groups A^0 ,
wherein:

A^0 is A^1 , A^2 or W^3 with the proviso that the conjugate includes at least one A^1 ;

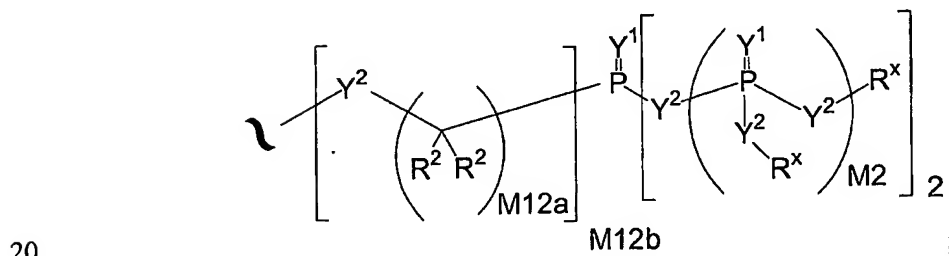
A^1 is:



A^2 is:



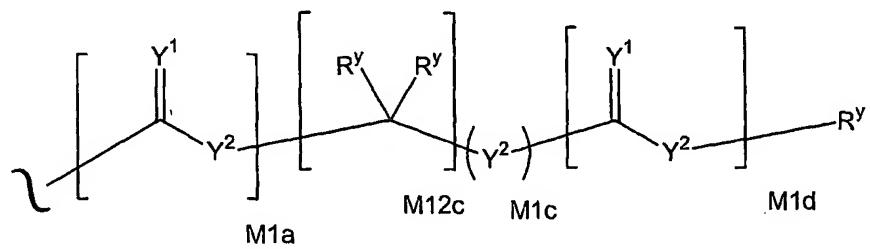
A^3 is:



Y^1 is independently O, S, $N(R^x)$, $N(O)(R^x)$, $N(OR^x)$, $N(O)(OR^x)$, or $N(N(R^x)(R^x))$;

Y^2 is independently a bond, O, $N(R^x)$, $N(O)(R^x)$, $N(OR^x)$, $N(O)(OR^x)$, $N(N(R^x)(R^x))$, $-S(O)_{M2-}$, or $-S(O)_{M2-}S(O)_{M2-}$; and when Y^2 joins two phosphorous atoms Y^2 can also be $C(R^2)(R^2)$;

R^x is independently H, R^1 , R^2 , W^3 , a protecting group, or the formula:



wherein:

R^y is independently H, W^3 , R^2 or a protecting group;

R^1 is independently H or alkyl of 1 to 18 carbon atoms;

R^2 is independently H, R^1 , R^3 or R^4 wherein each R^4 is independently substituted with 0 to 3 R^3 groups or taken together at a carbon atom, two R^2 groups form a ring of 3 to 8 carbons and the ring may be substituted with 0 to 3 R^3 groups;

R^3 is R^{3a} , R^{3b} , R^{3c} or R^{3d} , provided that when R^3 is bound to a heteroatom, then R^3 is R^{3c} or R^{3d} ;

R^{3a} is F, Cl, Br, I, -CN, N_3 or $-NO_2$;

R^{3b} is Y^1 ;

R^{3c} is $-R^x$, $-N(R^x)(R^x)$, $-SR^x$, $-S(O)R^x$, $-S(O)_2R^x$, $-S(O)(OR^x)$, $-S(O)_2(OR^x)$, $-OC(Y^1)R^x$, $-OC(Y^1)OR^x$, $-OC(Y^1)(N(R^x)(R^x))$, $-SC(Y^1)R^x$, $-SC(Y^1)OR^x$, $-SC(Y^1)(N(R^x)(R^x))$, $-N(R^x)C(Y^1)R^x$, $-N(R^x)C(Y^1)OR^x$, or $-N(R^x)C(Y^1)(N(R^x)(R^x))$;

R^{3d} is $-C(Y^1)R^x$, $-C(Y^1)OR^x$ or $-C(Y^1)(N(R^x)(R^x))$;

R^4 is an alkyl of 1 to 18 carbon atoms, alkenyl of 2 to 18 carbon atoms, or alkynyl of 2 to 18 carbon atoms;

R^5 is R^4 wherein each R^4 is substituted with 0 to 3 R^3 groups;

W^3 is W^4 or W^5 ;

W^4 is R^5 , $-C(Y^1)R^5$, $-C(Y^1)W^5$, $-SO_{M2}R^5$, or $-SO_{M2}W^5$;

W^5 is carbocycle or heterocycle wherein W^5 is independently substituted with 0 to 3 R^2 groups;

W^6 is W^3 independently substituted with 1, 2, or 3 A^3 groups;

$M2$ is 0, 1 or 2;

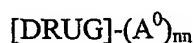
5 $M12a$ is 1, 2, 3, 4, 5, 6, 7, 8, 9, 10, 11 or 12;

$M12b$ is 0, 1, 2, 3, 4, 5, 6, 7, 8, 9, 10, 11 or 12;

$M1a$, $M1c$, and $M1d$ are independently 0 or 1; and

$M12c$ is 0, 1, 2, 3, 4, 5, 6, 7, 8, 9, 10, 11 or 12.

10 3. The conjugate of claim 2, or a pharmaceutically acceptable salt or solvate thereof, which has the formula:



wherein:

$DRUG$ is a compound of any one of formulae 500-547; and

15 m is 1, 2, or 3.

4. The conjugate of claim 2 which has any one of formulae 1-151 wherein:
one A^0 is A^1 ; each X^{50} is independently hydrogen, F, Cl, CF_3 , CN,
methyl, or *tert*-butyl;

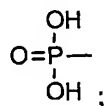
20 X^{51} is hydrogen, halo, trifluoromethyl, (C1-C3)alkyl, cyano, or
(C1-C3)alkoxy;

X^{52} is hydrogen, fluoro, chloro, bromo, methyl, or trifluoromethyl;

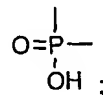
X^{53} is -O-, or -S-;

X^{54} and X^{55} are independently selected from hydrogen or a C_1 - C_{18} acyl;

25 X^{56} is hydrogen, a C_1 - C_{18} acyl, or



or X^{54} is hydrogen and together X^{55} and X^{56} are



X^{57} is H, amino, hydroxy, or a halogen selected from Cl and Br;

30 X^{58} is hydrogen, F, Cl, CF_3 , cyano, methyl, or *t*-butyl;

X^{59} is hydrogen, CH_2OH ;

X^{60} is $\text{CO}(\text{CH}_2)_6\text{CONMe}(\text{CH}_2)_2\text{SO}_3\text{H}$;

X^{62} is methyl, chloro, or trifluoromethyl;

X^{63} is H, methyl, ethyl, cyclopropyl, vinyl, or trifluoromethyl;

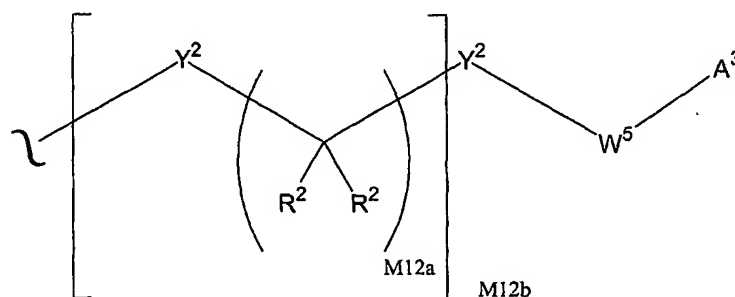
X^{64} is H, methyl, ethyl, cyclopropyl, chloro, vinyl, allyl, 3-methyl-1-

5 buten-1-yl;

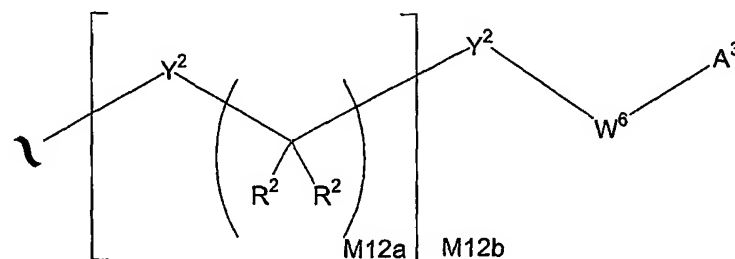
X^{65} is hydrogen or F; and

Ar is aryl or heteroaryl.

- 10 5. The conjugate of any one of claims 2-4, wherein each A^1 is of the formula:

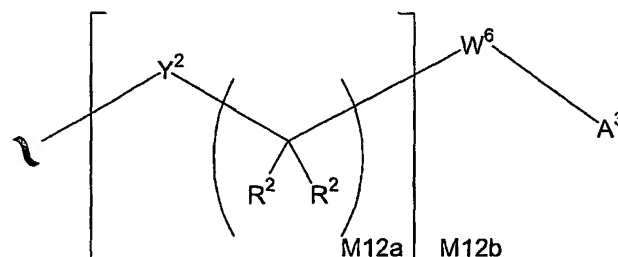


6. The conjugate of any one of claims 2-4, wherein each A^1 is of the formula:

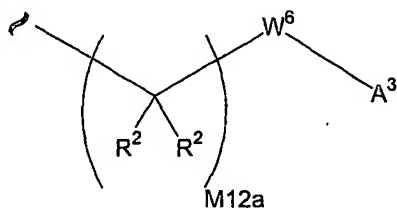


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7. The conjugate of any one of claims 2-4, wherein each A^1 is of the formula:

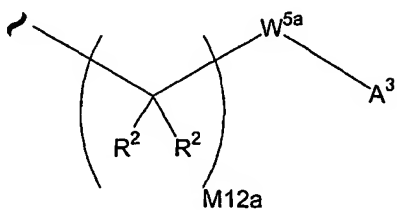


8. The conjugate of any one of claims 2-4, wherein each A^1 is of the formula:



5

9. The conjugate of any one of claims 2-4, wherein each A^1 is of the formula:



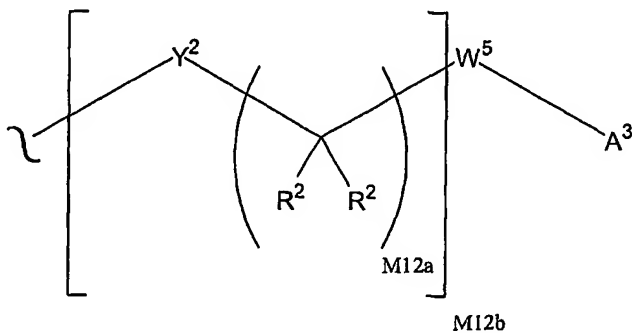
10

wherein: W^{5a} is a carbocycle or a heterocycle where W^{5a} is independently substituted with 0 or 1 R^2 groups.

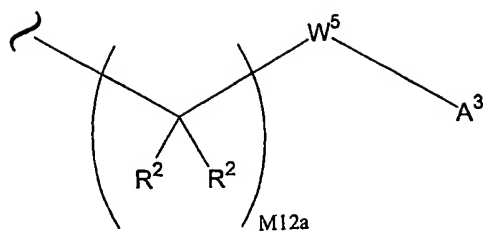
10. The conjugate of any one of claims 2-4, wherein M12a is 1.

15

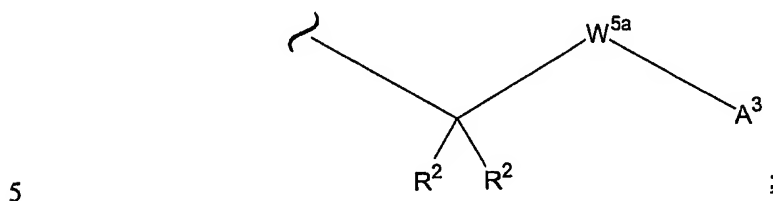
11. The conjugate of any one of claims 2-4, wherein each A^1 is of the formula:



- 20 12. The conjugate of any one of claims 2-4, wherein each A^1 is of the formula:

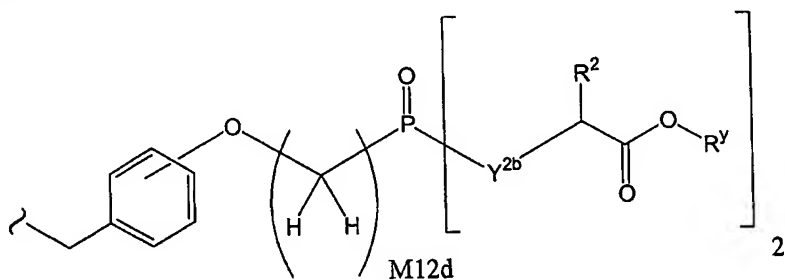


13. The conjugate of any one of claims 2-4, wherein each A¹ is of the formula:



wherein: W^{5a} is a carbocycle independently substituted with 0 or 1 R² groups.

14. The conjugate of any one of claims 2-4, wherein each A¹ is of the formula:

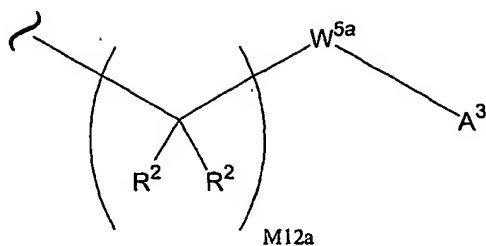


wherein:

Y^{2b} is O or N(R²); and

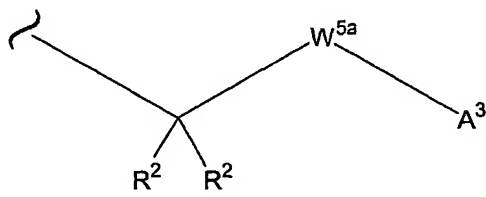
M12d is 1, 2, 3, 4, 5, 6, 7 or 8.

15. The conjugate of any one of claims 2-4, wherein each A¹ is of the formula:



wherein: W^{5a} is a carbocycle independently substituted with 0 or 1 R^2 groups.

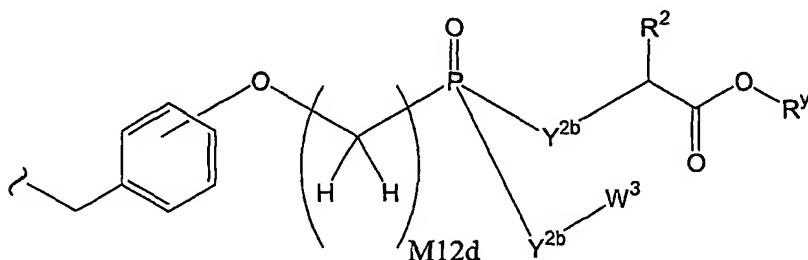
16. The conjugate of any one of claims 2-4, wherein each A^1 is of the
5 formula:



wherein: W^{5a} is a carbocycle or heterocycle where W^{5a} is independently substituted with 0 or 1 R^2 groups.

10

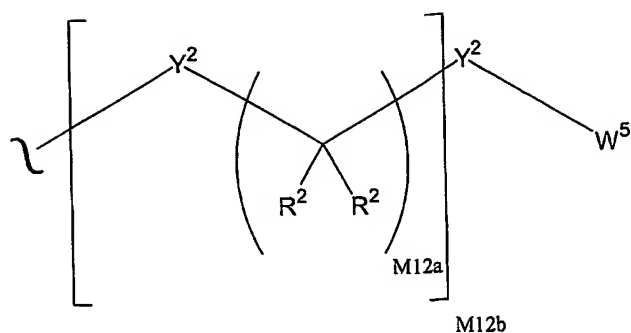
17. The conjugate of any one of claims 2-4, wherein each A^1 is of the
formula:



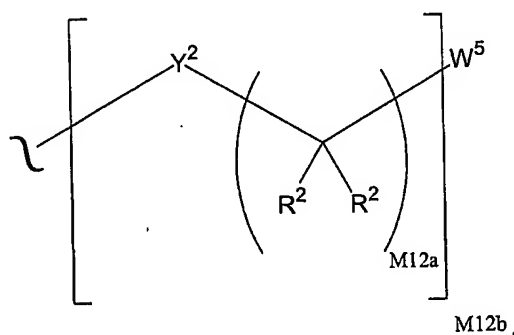
wherein:

- 15 Y^{2b} is O or $N(R^2)$; and
 M12d is 1, 2, 3, 4, 5, 6, 7 or 8.

18. The conjugate of any one of claims 2-17 wherein each A^2 is of the
formula:



19. The conjugate of any one of claims 2-17 wherein each A^2 is of the formula:



5

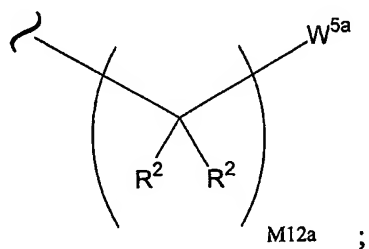
20. The conjugate of any one of claims 2-17 wherein each M12b is 1.

21. The conjugate of claim 20 where M12b is 0, Y^2 is a bond and W^5 is a carbocycle or heterocycle where W^5 is optionally and independently substituted with 1, 2, or 3 R^2 groups.

10

22. The conjugate of any one of claims 2-17 wherein each A^2 is of the formula:

15



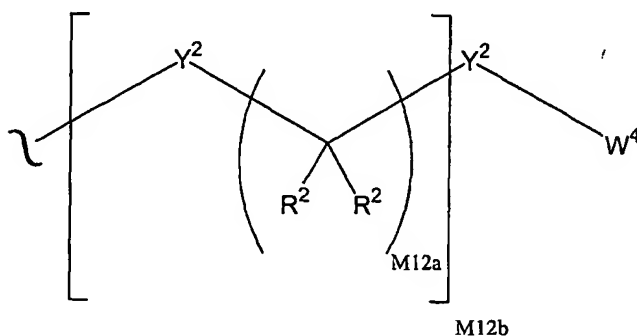
wherein: W^{5a} is a carbocycle or heterocycle where W^{5a} is optionally and independently substituted with 1, 2, or 3 R^2 groups.

23. The conjugate of claim 22 wherein M12a is 1.

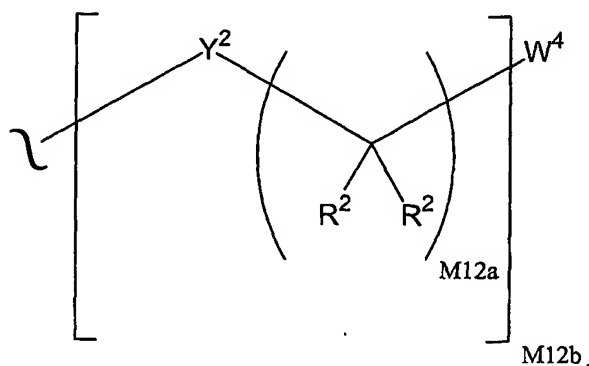
5

24. The conjugate of any one of claims 2-17 wherein each A^2 is selected from phenyl, substituted phenyl, benzyl, substituted benzyl, pyridyl and substituted pyridyl.

10 25. The conjugate of any one of claims 2-17 wherein each A^2 is of the formula:

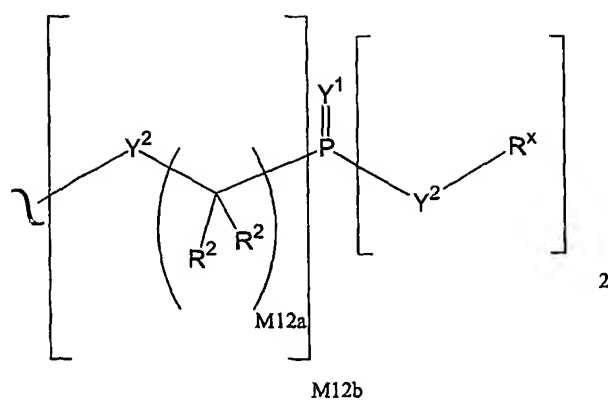


15 26. The conjugate of any one of claims 2-17 wherein each A^2 is of the formula:

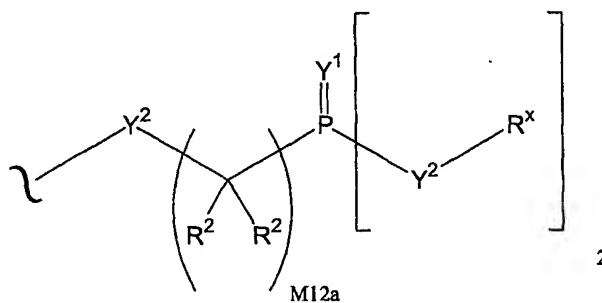


27. The conjugate of claim 26 wherein M12b is 1.

20 28. The conjugate of any one of claims 2-27 wherein each A^3 is of the formula:

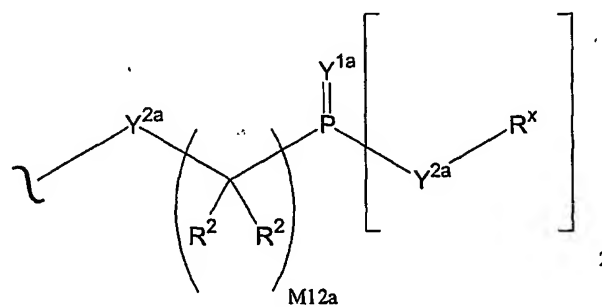


29. The conjugate of any one of claims 2-27 wherein each A³ is of the formula:



5

30. The conjugate of any one of claims 2-27 wherein each A³ is of the formula:



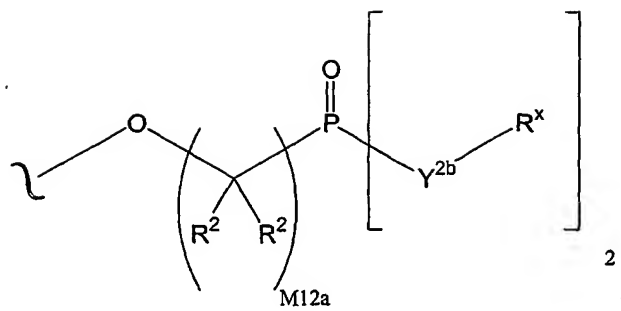
10 wherein:

Y^{1a} is O or S; and

Y^{2a} is O, N(R^x) or S.

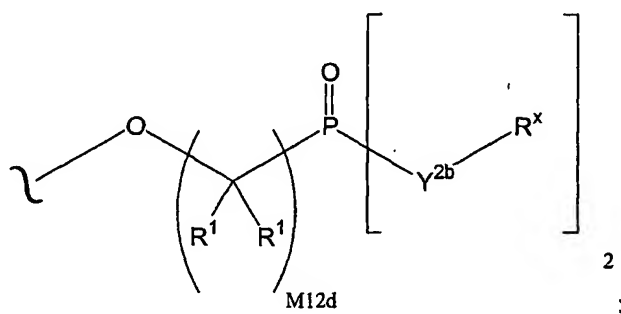
31. The conjugate of any one of claims 2-27 wherein each A³ is of the formula:

15



wherein Y^{2b} is O or $N(R^x)$.

32. The conjugate of any one of claims 2-27 wherein each A^3 is of the
5 formula:



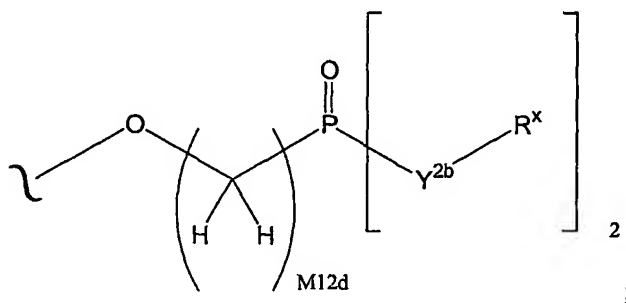
wherein:

Y^{2b} is O or $N(R^x)$; and

M12d is 1, 2, 3, 4, 5, 6, 7 or 8.

10

33. The conjugate of any one of claims 2-27 wherein each A^3 is of the
formula:



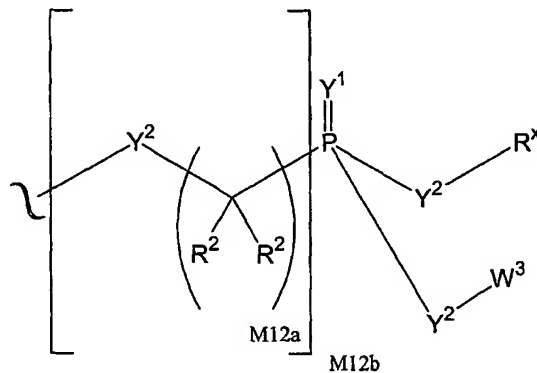
wherein:

15 Y^{2b} is O or $N(R^x)$; and

M12d is 1, 2, 3, 4, 5, 6, 7 or 8.

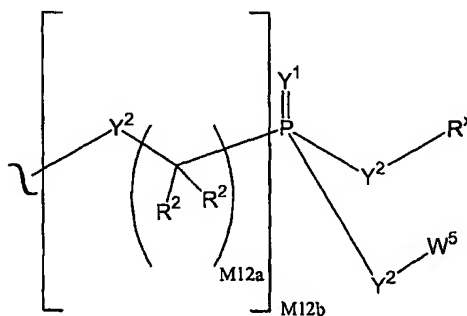
34. The conjugate of claim 33 wherein M12d is 1.

35. The conjugate of any one of claims 2-27 wherein each A³ is of the formula:



5

36. The conjugate of any one of claims 2-27 wherein each A³ is of the formula:

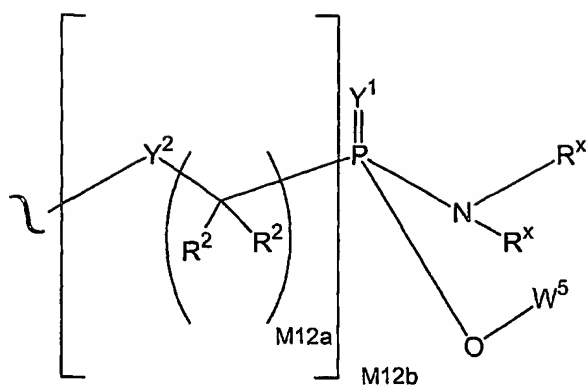


10

37. The conjugate of claim 36 wherein W⁵ is a carbocycle.

38. The conjugate of any one of claims 2-27 wherein each A³ is of the formula:

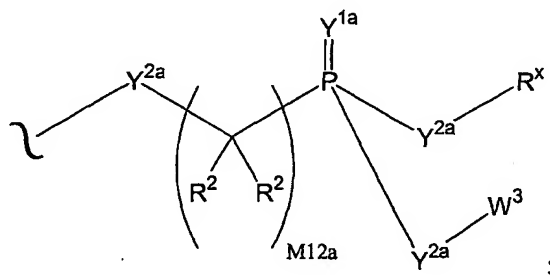
15



39. The conjugate of claim 38 wherein W⁵ is phenyl.

5 40. The conjugate of claim 39 wherein M12b is 1.

41. The conjugate of any one of claims 2-27 wherein each A³ is of the formula:



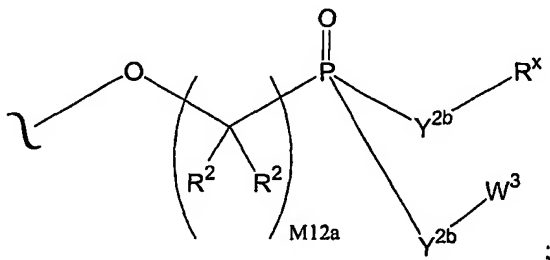
10 wherein:

Y^{1a} is O or S; and

Y^{2a} is O, N(R^x) or S.

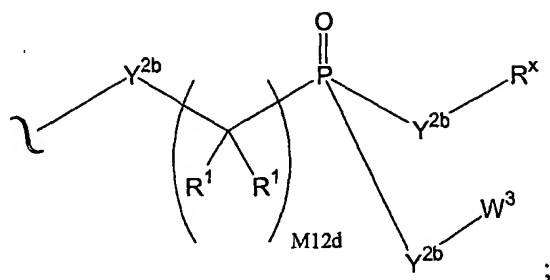
42. The conjugate of any one of claims 2-27 wherein each A³ is of the

15 formula:



wherein Y^{2b} is O or N(R^x).

43. The conjugate of any one of claims 2-27 wherein each A^3 is of the formula:



5 wherein:

Y^{2b} is O or $N(R^x)$; and

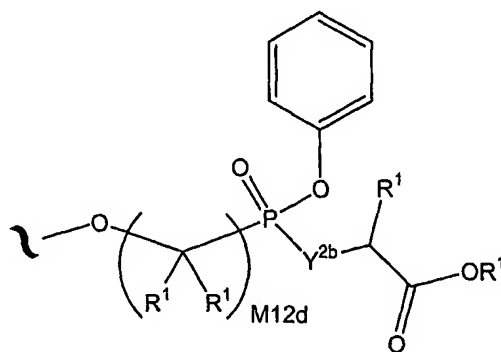
M12d is 1, 2, 3, 4, 5, 6, 7 or 8.

44. The conjugate of claim 43 wherein R^1 is H.

10

45. The conjugate of claim 44 wherein M12d is 1.

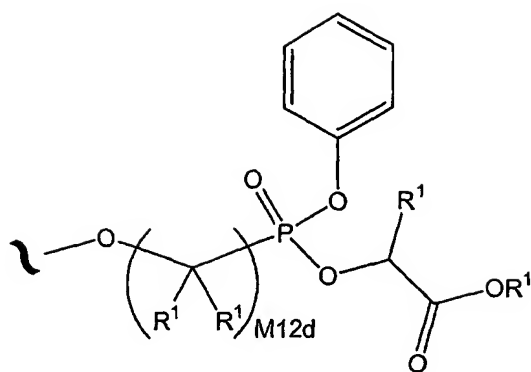
46. The conjugate of any one of claims 2-27 wherein each A^3 is of the formula:



15

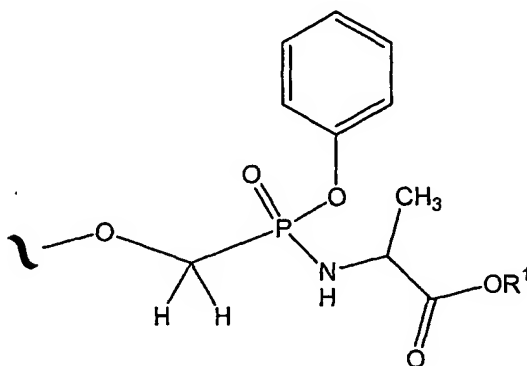
wherein the phenyl carbocycle is substituted with 0, 1, 2, or 3 R^2 groups.

47. The conjugate of any one of claims 2-27 wherein each A^3 is of the formula:

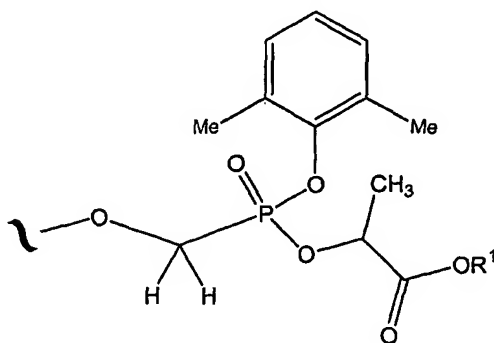


wherein the phenyl carbocycle is substituted with 0, 1, 2, or 3 R^2 groups.

48. The conjugate of any one of claims 2-27 wherein each A^3 is of the
5 formula:

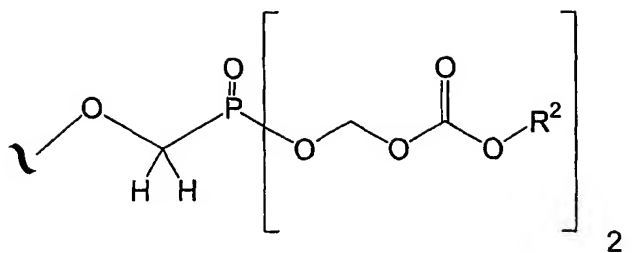


49. The conjugate of any one of claims 2-27 wherein each A^3 is of the
formula:

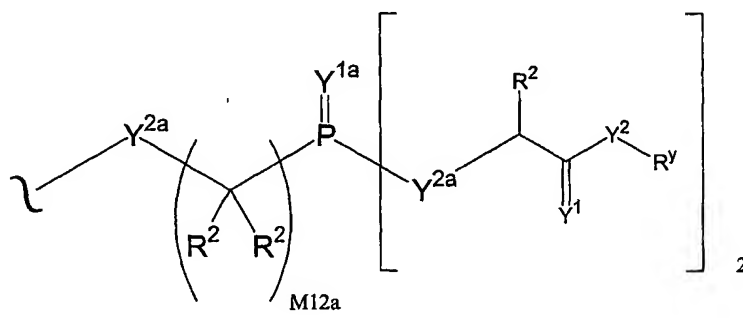


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50. The conjugate of any one of claims 2-27 wherein each A^3 is of the
formula:



51. The conjugate of any one of claims 2-27 wherein each A^3 is of the formula:

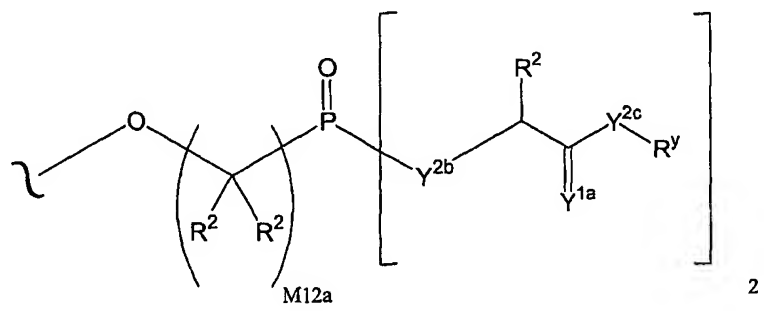


wherein:

Y^{1a} is O or S; and

Y^{2a} is O, $N(R^2)$ or S.

52. The conjugate of any one of claims 2-27 wherein each A^3 is of the formula:



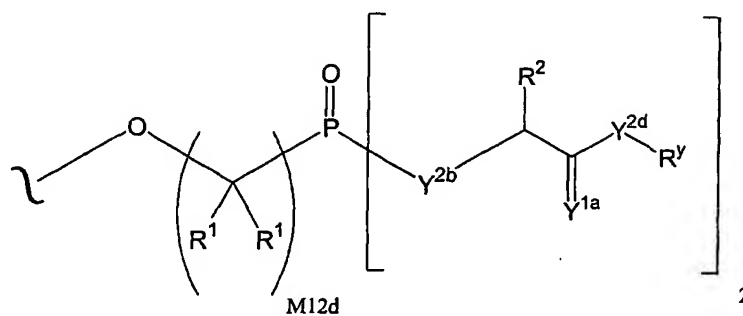
wherein:

Y^{1a} is O or S;

Y^{2b} is O or $N(R^2)$; and

Y^{2c} is O, $N(R^y)$ or S.

53. The conjugate of any one of claims 2-27 wherein each A^3 is of the formula:



5 wherein:

Y^{1a} is O or S;

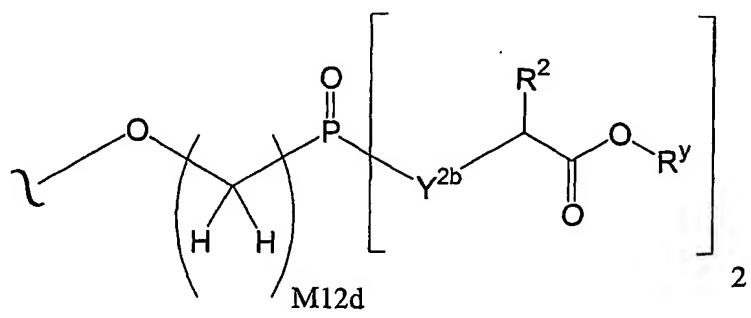
Y^{2b} is O or $N(R^2)$;

Y^{2d} is O or $N(R^y)$; and

M12d is 1, 2, 3, 4, 5, 6, 7 or 8.

10

54. The conjugate of any one of claims 2-27 wherein each A^3 is of the formula:



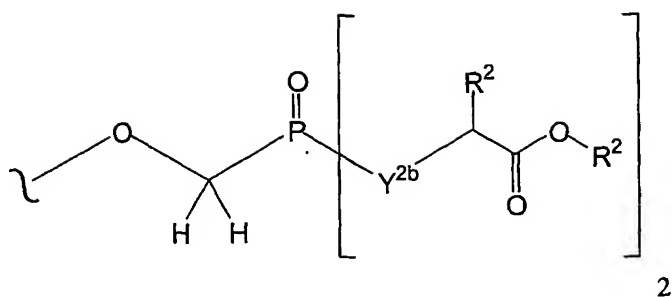
15 wherein:

Y^{2b} is O or $N(R^2)$; and

M12d is 1, 2, 3, 4, 5, 6, 7 or 8.

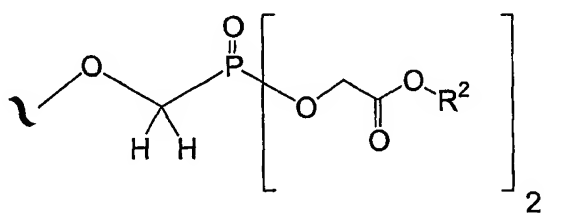
55. The conjugate of any one of claims 2-27 wherein each A^3 is of the formula:

20

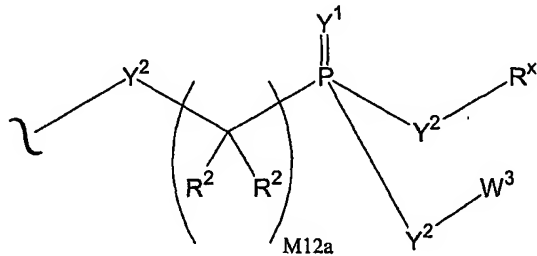


wherein Y^{2b} is O or $\text{N}(\text{R}^2)$.

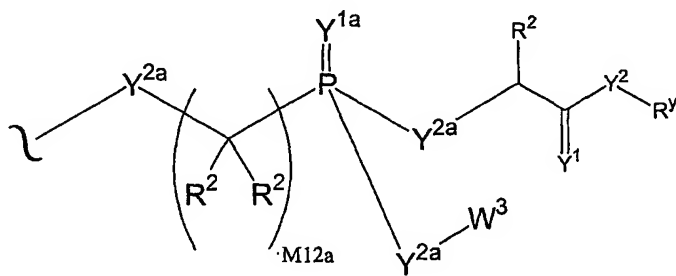
56. The conjugate of any one of claims 2-27 wherein each A^3 is of the
5 formula:



57. The conjugate of any one of claims 2-27 wherein each A^3 is of the
formula:



58. The conjugate of any one of claims 2-27 wherein each A^3 is of the
formula:

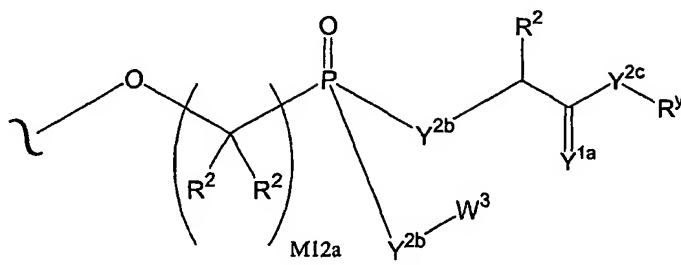


wherein:

Y^{1a} is O or S; and

Y^{2a} is O, N(R²) or S.

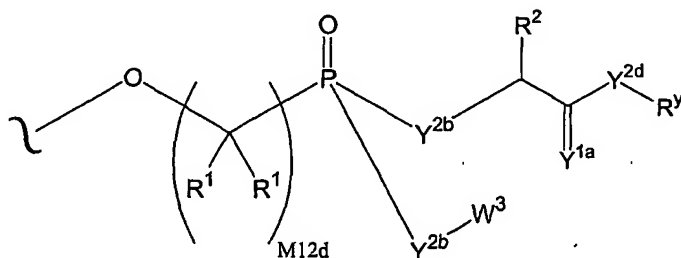
- 5 59. The conjugate of any one of claims 2-27 wherein each A³ is of the formula:



wherein:

- 10 Y^{1a} is O or S;
 Y^{2b} is O or N(R²); and
 Y^{2c} is O, N(R^y) or S.

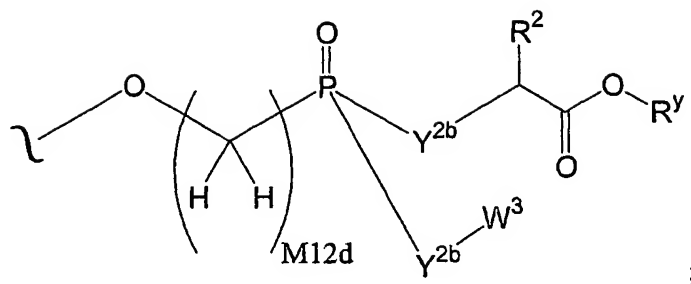
60. The conjugate of any one of claims 2-27 wherein each A³ is of the
 15 formula:



wherein:

- 20 Y^{1a} is O or S;
 Y^{2b} is O or N(R²);
 Y^{2d} is O or N(R^y); and
 M12d is 1, 2, 3, 4, 5, 6, 7 or 8.

61. The conjugate of any one of claims 2-27 wherein each A³ is of the
 25 formula:



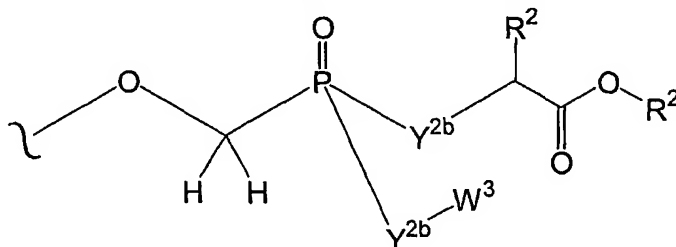
wherein:

Y^{2b} is O or $N(R^2)$; and

M12d is 1, 2, 3, 4, 5, 6, 7 or 8.

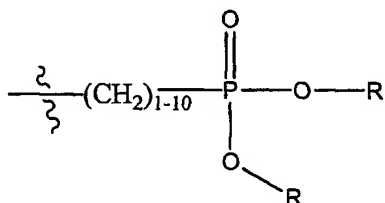
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62. The conjugate of any one of claims 2-27 wherein each A^3 is of the formula:



wherein Y^{2b} is O or $N(R^2)$.

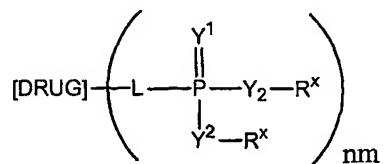
10 63. The conjugate of claim 3 wherein A^0 is of the formula:



wherein each R is independently alkyl.

15

64. The conjugate of claim 1, 2, 3, or 4 which has the formula:



or a pharmaceutically acceptable salt or solvate thereof,

wherein:

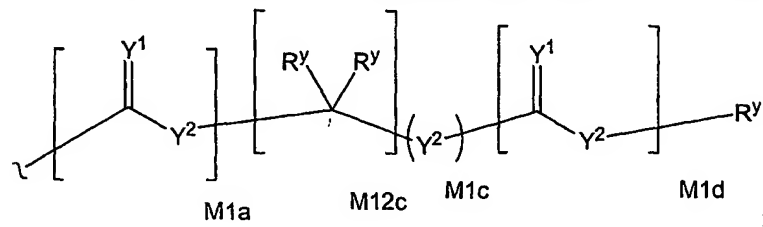
DRUG is an with immuno-modulatory compound;

Y^1 is independently O, S, $N(R^x)$, $N(O)(R^x)$, $N(OR^x)$, $N(O)(OR^x)$, or

5 $N(N(R^x)(R^x))$;

Y^2 is independently a bond, O, $N(R^x)$, $N(O)(R^x)$, $N(OR^x)$, $N(O)(OR^x)$, $N(N(R^x)(R^x))$, $-S(O)_{M2}$, or $-S(O)_{M2}-S(O)_{M2}$;

R^x is independently H, R^2 , W^3 , a protecting group, or the formula:



10 R^y is independently H, W^3 , R^2 or a protecting group;

R^2 is independently H, R^3 or R^4 wherein each R^4 is independently substituted with 0 to 3 R^3 groups;

R^3 is R^{3a} , R^{3b} , R^{3c} or R^{3d} , provided that when R^3 is bound to a heteroatom, then R^3 is R^{3c} or R^{3d} ;

15 R^{3a} is F, Cl, Br, I, -CN, N_3 or $-NO_2$;

R^{3b} is Y^1 ;

R^{3c} is $-R^x$, $-N(R^x)(R^x)$, $-SR^x$, $-S(O)R^x$, $-S(O)_2R^x$, $-S(O)(OR^x)$, $-S(O)_2(OR^x)$, $-OC(Y^1)R^x$, $-OC(Y^1)OR^x$, $-OC(Y^1)(N(R^x)(R^x))$, $-SC(Y^1)R^x$, $-SC(Y^1)OR^x$, $-SC(Y^1)(N(R^x)(R^x))$, $-N(R^x)C(Y^1)R^x$, $-N(R^x)C(Y^1)OR^x$, or $-N(R^x)C(Y^1)(N(R^x)(R^x))$;

R^{3d} is $-C(Y^1)R^x$, $-C(Y^1)OR^x$ or $-C(Y^1)(N(R^x)(R^x))$;

R^4 is an alkyl of 1 to 18 carbon atoms, alkenyl of 2 to 18 carbon atoms, or alkynyl of 2 to 18 carbon atoms;

25 R^5 is R^4 wherein each R^4 is substituted with 0 to 3 R^3 groups;

W^3 is W^4 or W^5 ;

W^4 is R^5 , $-C(Y^1)R^5$, $-C(Y^1)W^5$, $-SO_2R^5$, or $-SO_2W^5$;

W^5 is carbocycle or heterocycle wherein W^5 is independently substituted with 0 to 3 R^2 groups;

30 $M2$ is 1, 2, or 3;

M1a, M1c, and M1d are independently 0 or 1;

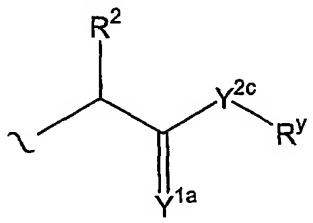
M12c is 0, 1, 2, 3, 4, 5, 6, 7, 8, 9, 10, 11 or 12;

nn is 1, 2, or 3; and

L is a direct bond or a linking group.

5

65. The conjugate of claim 64 wherein each R^x is of the formula:

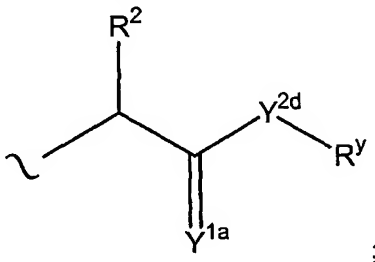


Y^{1a} is O or S; and

Y^{2c} is O, N(R^y) or S.

10

66. The conjugate of claim 64 wherein each R^x is of the formula:

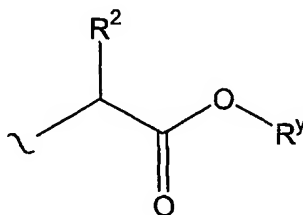


Y^{1a} is O or S; and

Y^{2d} is O or N(R^y).

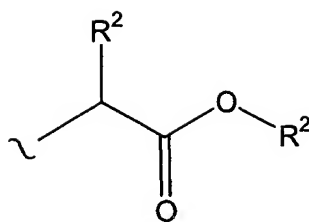
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67. The conjugate of claim 64 wherein each R^x is of the formula:

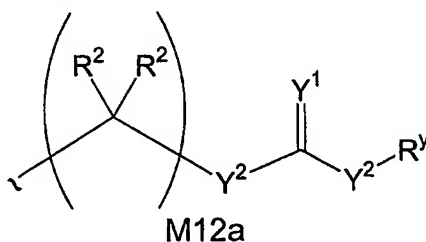


68. The conjugate of any one of claims 65-67 wherein each R^y is
20 independently H or alkyl of 1 to 10 carbons.

69. The conjugate of claim 64 wherein each R^x is of the formula:

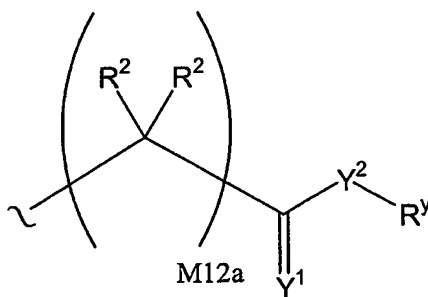


70. The conjugate of claim 64 wherein each R^x is of the formula:



5

71. The conjugate of claim 64 wherein each R^x is of the formula:



72. The conjugate of claim 64 wherein each Y^1 is O or S.

10

73. The conjugate of claim 64 wherein each Y^2 is O, $N(R^y)$ or S.

74. The conjugate of any one of claims 64-73 wherein nn is 1.

- 15 75. The conjugate of any one of claims 64-73 wherein nn is 2.

76. The conjugate of any one of claims 64-73 wherein nn is 3.

- 20 77. The conjugate of claim 64 wherein the kinase inhibiting compound is a compound of any one of formulae 500-547.

78. The conjugate of claim 77 wherein each L has a molecular weight of from about 20 daltons to about 400 daltons.
- 5 79. The conjugate of claim 77 wherein each L has a length of about 5 angstroms to about 300 angstroms.
80. The conjugate of claim 77 wherein each L separates the compound of any one of formulae 500-547 and the phosphorous of the phosphonate group by
10 about 5 angstroms to about 200 angstroms, inclusive.
81. The conjugate of claim 77 wherein each L is a divalent, branched or unbranched, saturated or unsaturated, hydrocarbon chain, having from 2 to 25 carbon atoms, wherein one or more of the carbon atoms is optionally replaced by
15 (-O-), and wherein the chain is optionally substituted on carbon with one or more substituents selected from (C₁-C₆)alkoxy, (C₃-C₆)cycloalkyl, (C₁-C₆)alkanoyl, (C₁-C₆)alkanoyloxy, (C₁-C₆)alkoxycarbonyl, (C₁-C₆)alkylthio, azido, cyano, nitro, halo, hydroxy, oxo (=O), carboxy, aryl, aryloxy, heteroaryl, and heteroaryloxy.
- 20 82. The conjugate of claim 77 wherein each L is of the formula W-A wherein A is (C₁-C₂₄)alkylene, (C₂-C₂₄)alkenylene, (C₂-C₂₄)alkynylene, (C₃-C₈)cycloalkylene, (C₆-C₁₀)aryl or a combination thereof, wherein each W is -N(R)C(=O)-, -C(=O)N(R)-, -OC(=O)-, -C(=O)O-, -O-, -S-, -S(O)-, -S(O)₂-,
25 -N(R)-, -C(=O)-, -N(R)C=N(R)-N(R)-, -C(R)=N(R)-, -S(O)_{M2}-N(R)-, -N(R)-S(O)_{M2}-, or a direct bond; wherein each R is independently H or alkyl of 1 to 10 carbons.
83. The conjugate of claim 82 wherein each A is alkylene of 1 to 10 carbons.
- 30 84. The conjugate of claim 77 wherein each L is a divalent radical formed from a peptide.

85. The conjugate of claim 77 wherein each L is a divalent radical formed from an amino acid.
86. The conjugate of claim 77 wherein each L is a divalent radical formed from poly-L-glutamic acid, poly-L-aspartic acid, poly-L-histidine, poly-L-ornithine, poly-L-serine, poly-L-threonine, poly-L-tyrosine, poly-L-leucine, poly-L-lysine-L-phenylalanine, poly-L-lysine or poly-L-lysine-L-tyrosine.
87. The conjugate of claim 77 wherein each L is of the formula $W-(CH_2)_n$ wherein, n is between about 1 and about 10; and W is $-N(R)C(=O)-$, $-C(=O)N(R)-$, $-OC(=O)-$, $-C(=O)O-$, $-O-$, $-S-$, $-S(O)-$, $-S(O)_2-$, $-C(=O)-$, $-N(R)-$, $-N(R)C=N(R)-N(R)-$, $-C(R)=N(R)-$, $-S(O)_{M2}-N(R)-$, $-N(R)-S(O)_{M2}-$, or a direct bond; wherein each R is independently H or (C_1-C_6) alkyl.
88. The conjugate of claim 77 wherein each L is methylene, ethylene, or propylene.
89. The conjugate of claim 77 wherein each L is linked to P at a carbon atom of L.
90. The conjugate as described in any one of claims 1-89, which is isolated and purified.
91. The conjugate of any one of claims 1-90 which is not an anti-inflammatory compound.
92. The conjugate of any one of claims 1-91 which is not an anti-infective.
93. The conjugate of any one of claims 1-92 which is not a compound that is not a kinase inhibitor.
94. The conjugate of any one of claims 1-93 which is not a compound that is active against metabolic diseases.

95. The conjugate of any one of claims 1-94 which is not an antiviral agent.
96. The conjugate of any one of claims 1-95 which is not a nucleoside.
- 5 97. The conjugate of any one of claims 1-96 which is not a IMPDH inhibitor.
98. The conjugate of any one of claims 1-97 which is not an antimetabolite.
- 10 99. The conjugate of any one of claims 1-98 which is not a PNP inhibitor.
100. The conjugate of any one of claims 2-99 which is not a substituted compound of any one of formulae 500-533, 535-541, or 543-547.
- 15 101. The conjugate of any one of claims 2-100 which is not a substituted compound of formula 534 or 542.
102. The conjugate of any one of claims 4-101 which is not a compound of any one of formulae 1-104, 107-124, or 128-151.
- 20 103. The conjugate of any one of claims 4-102 which is not a compound of formula 105-106, or 125-127.
104. A kinase inhibitor conjugate as described herein.
- 25 105. A compound of the formula MBF.
106. A compound of claim 105 selected from Table 100.
- 30 107. A pharmaceutical composition comprising a pharmaceutically acceptable excipient and a conjugate as described in any one of claims 1-89 and 91-104 or a compound as described in claim 105 or 106.

108. A unit dosage form comprising a conjugate as described in any one of claims 1-89 and 91-104 or a compound as described in claim 105 or 106 and a pharmaceutically acceptable excipient.
- 5 109. A method for inhibiting an immune response *in vitro* or *in vivo* comprising contacting a sample in need of such treatment with a conjugate as described in any one of claims 1-89 and 91-104 or a compound as described in claim 105 or 106.
- 10 110. The method of claim 109 wherein the contacting is *in vivo*.
111. The invention also provides a method of modulating an immune response in a mammal, comprising administering a compound as described in any one of claims 1-89 and 91-104 or a compound as described in claim 105 or 106 to the
15 mammal.
112. The method of claim 111 wherein the compound is formulated with a pharmaceutically acceptable carrier.
- 20 113. The method of claim 112 wherein the formulation further comprises a second active ingredient.
114. The method of claim 111 or 112 wherein an immune response is suppressed.
25
115. A conjugate as described in any one of claims 1-89 and 91-104 or a compound as described in claim 105 or 106 for use in medical therapy.
116. The use of a conjugate as described in any one of claims 1-89 and 91-104
30 or a compound as described in claim 105 or 106 to prepare a medicament for modulating an immune response in an animal.
117. A phosphonate substituted immuno-modulatory conjugate as described herein.

118. A method for preparing a conjugate as described in the schemes and examples herein.